

POINT SOURCE MODEL
EVALUATION AND DEVELOPMENT
STUDY

Appendix C - User Guide to IMPACT

prepared by

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California Air Resources Board
and the
California Energy Resources Conservation
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USER GUIDE TO IMPACT
INTEGRATED MODEL FOR PLUMES AND ATMOSPHERICS
IN COMPLEX TERRAIN

APPENDIX C

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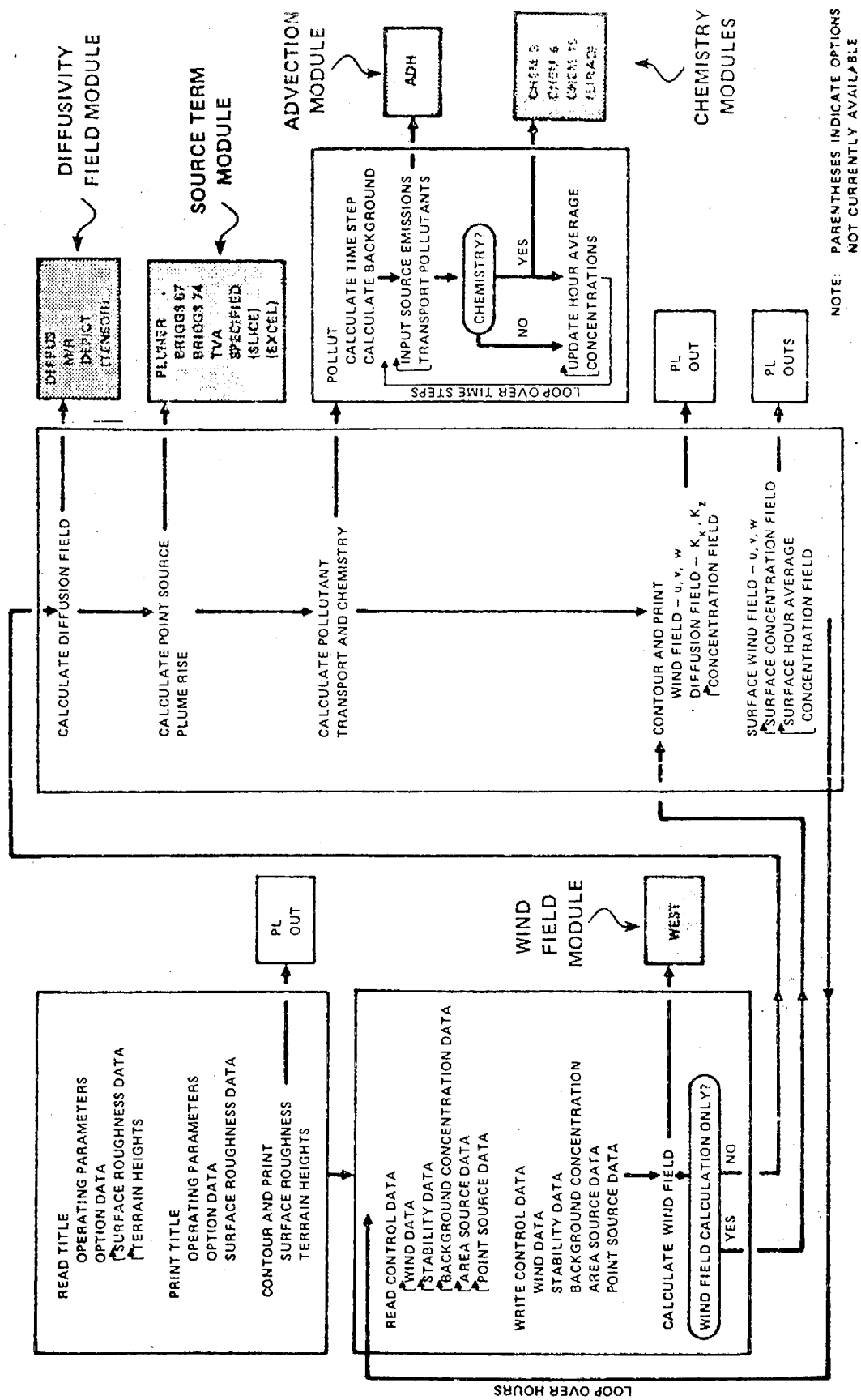
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1. INTRODUCTION

The Integrated Model for Plumes and Atmospherics in Complex Terrain (IMPACT) is the computer code for a three-dimensional grid model for calculating the impact of pollutants, either inert or reactive, in simple or complex terrain, emitted from either point or area sources. As such, it is one of the most comprehensive air pollution model yet developed. It automatically treats single or multiple point or area sources, the effects of arbitrary vertical temperature stratifications (multiple inversions) on the wind and diffusion fields, shear flows caused by the atmospheric boundary layer or by terrain effects, and chemical transformations (such as those creating photochemical smog).

The code was written in modular format such that the incorporation of new or alternative formulations for plume rise, winds diffusion, advection, and chemistry is particularly straightforward with minimal changes required in the existing program. A macro-flow diagram of IMPACT is shown in Figure 1. The shaded boxes represent the five modules that are the major components of IMPACT; the rest of the code is concerned with program control and input and output functions.

The next section of this user guide contains a description of the various components of the IMPACT code. These descriptions may contain flow diagrams, input/output variables, equations and graphs as required to describe the routines. Section 3 presents a discussion of the various options available in the code, including selection of chemistries, diffusion models, averaging duration, and output plots and edits. Also included in this section is a discussion of the limitations on problem size, data input, etc. A detailed discussion of the input data and source file structure for IMPACT is presented in section 4. Also included is a discussion on how to modify the common storage block for different size problems. Section 5 provides helpful caveats for IMPACT usage. A sample run concludes the document.



NOTE: PARENTHESES INDICATE OPTIONS
NOT CURRENTLY AVAILABLE

Figure 1. Macro-Flow Diagram of IMPACT.

IMPACT is written in FORTRAN IV and should be transferable to any computer system supporting a FORTRAN IV compiler. It is currently running on IBM 370/108 and DEC-10 systems.

2. PROGRAM ROUTINES

The IMPACT code is designed to solve the following conservation of mass equation:

$$\begin{aligned}
 \frac{\partial C_i}{\partial t} = & - \underbrace{\frac{\partial u C_i}{\partial x} - \frac{\partial v C_i}{\partial y} - \frac{\partial w C_i}{\partial z}}_{\text{advection}} \\
 & + \underbrace{\frac{\partial}{\partial x} \left(K_x \frac{\partial C_i}{\partial x} \right) + \frac{\partial}{\partial y} \left(K_y \frac{\partial C_i}{\partial y} \right) + \frac{\partial}{\partial z} \left(K_z \frac{\partial C_i}{\partial z} \right)}_{\text{diffusion}} \\
 & + \underbrace{S_i}_{\text{source term}} + \underbrace{\alpha_{ij} C_j + \beta_{ijk} C_i C_j}_{\text{chemistry}}
 \end{aligned}$$

where

- C_i = the concentration of the i^{th} pollution species
- u, v, w = the three-dimensional flow field vectors
- K_x, K_y, K_z = the three-dimensional eddy diffusivity
- S_i = the source term for the i^{th} pollutant species
- α, β = the chemical rate constants for the first and second order chemical reactions.

A flow diagram, by subroutine, for the IMPACT code is shown in Figure 2. The shaded subroutines represent the major components of the code; the remainder of routines direct program flow and process input and output data. A brief definition and a description of the inputs and outputs for each subroutine in IMPACT is listed in Table 1. A description of the program common block, the five major subroutines, and the input/output routines conclude this section.

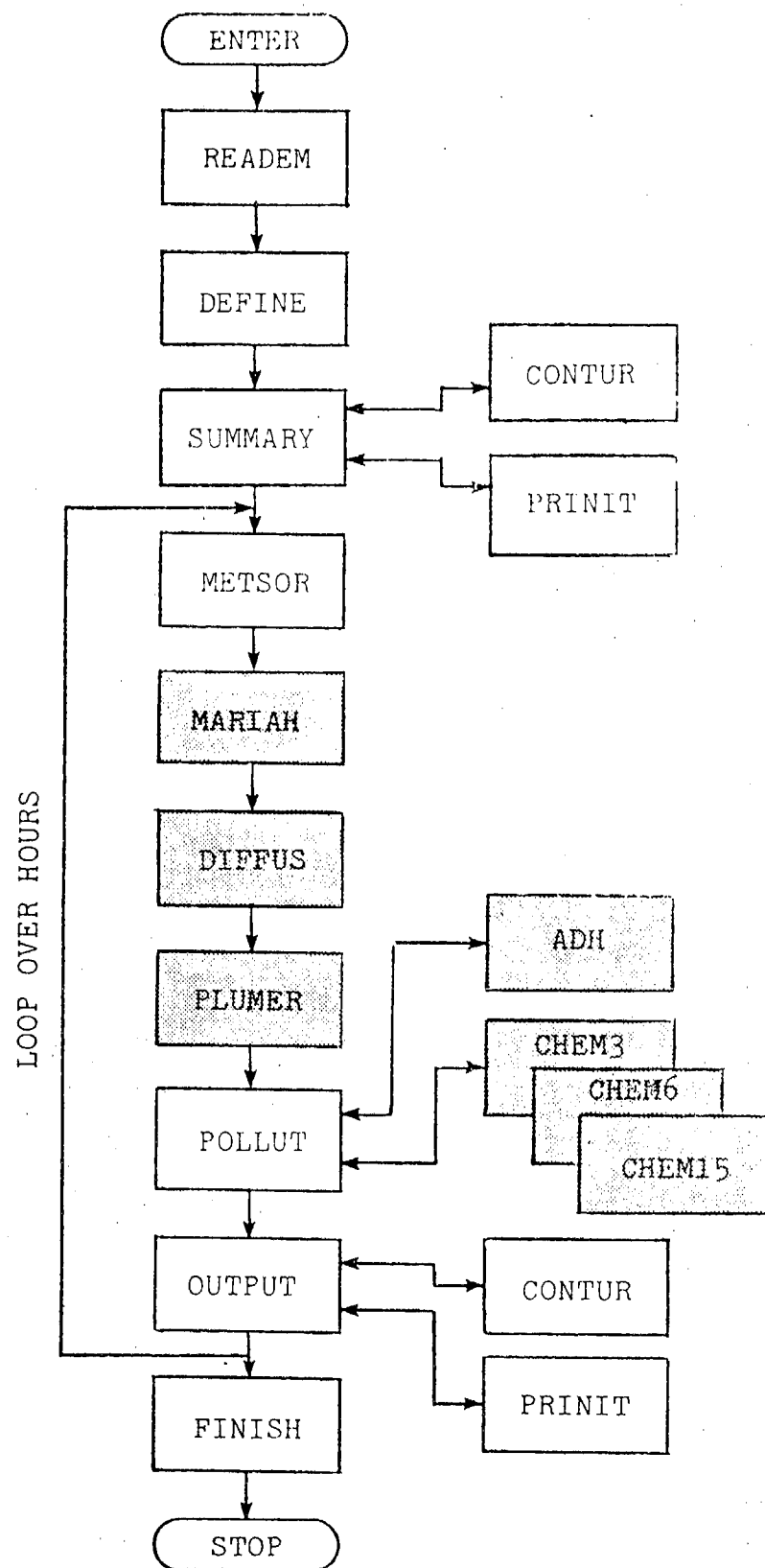


Figure 2. IMPACT Flow Chart.

Table 1. Description of Impact Subroutines.

SUBROUTINE	DESCRIPTION	INPUTS	OUTPUTS
READEM	READ SPACE, TIME INDEPENDENT QUANTITIES, OPTION, ETC.	NONE	GRID SIZE, OPTIONS, ETC.
DEFINE	ESTABLISH USEFUL INVARIANT QUANTITIES FOR LATER PROGRAM EFFICIENCY	GRID SIZE, CELL SIZE, ETC.	$1/\Delta x$, $1/\Delta x^2$, etc.
SUMMARY	PRINT A SUMMARY OF THE INITIAL STATE OF THE SYSTEM; READ TERRAIN AND SURFACE ROUGHNESS DATA	TERRAIN DATA, SURFACE ROUGHNESS DATA, PROGRAM INVARIANTS	INTERGERIZED TERRAIN ARRAY, INTERPOLATED SURFACE ROUGHNESS ARRAY, PRINTS AND EDITS OF INITIAL DATA
CONTUR	PRINT CONTOUR PLOTS FROM ANY SUBMITTED 2-D OR 3-D ARRAY	ANY 2-D OR 3-D ARRAY	CONTOUR PLOTS
PRINTIT	PRINTS THE FIELD ARRAYS TO THE FIRST TWO SIGNIFICANT DIGITS FROM ANY SUBMITTED 2-D OR 3-D ARRAY	ANY 2-D or 3-D ARRAY	PRINTER EDITS
METSOR	HOURLY READS THE METEOROLOGI- CAL SOURCE AND BACKGROUND DATA. PRINTS SUMMARY OF INPUT DATA. CONVERTS INPUT EMISSIONS FROM G/SEC TO PPM/SEC	WIND, STABILITY, POINT SOURCE, AREA SOURCE, BACKGROUND DATA	NET, SOURCE AND BACKGROUND DATA
MARIAH	CONTAINS THE SUBPROGRAM WESF. INTERPOLATES WIND, STABILITY DATA; CREATES DIVERGENCE-FREE WIND FIELD, CONSISTENT WITH WIND DATA, STABILITY DATA & TERRAIN	WIND AND STABILITY DATA, TERRAIN DATA	CONSISTENT WIND AND STABILITY ARRAYS
DIFFUS	CREATES A DIFFUSIVITY FIELD BASED ON EITHER THE M/R OR DEPICT MODELS	WIND AND STABILITY FIELDS, TERRAIN AND SURFACE ROUGHNESS DATA	DIFFUSIVITY FIELDS
PLUMER	DETERMINES THE POINT SOURCE PLUME RISE BASED ON BRIGGS OR FVA MODELS OR ON USER-SPECIFIED DATA	WIND AND STABILITY FIELDS, TERRAIN AND POINT SOURCE DATA	POSITION OF THE HORIZONTAL PLUME

Table 1. Description of Impact Subroutines (Continued).

SUBROUTINE	DESCRIPTION	INPUTS	OUTPUTS
POLLUT	CALCULATES POLLUTANT TRANSPORT, DIFFUSION, SOURCES AND CHEMISTRY CALCULATES AVERAGE SURFACE CONCENTRATION	WIND AND DIFFUSIVITY FIELDS, EMISSIONS, BACKGROUND CONCENTRATIONS, POSITION OF ELEVATED FLUXES	POLLUTANT CONCENTRATIONS IN TIME AND SPACE
ADL	UPDATES POLLUTANT CONCENTRATIONS DUE TO ADVECTION AND DIFFUSION USING A SECOND-ORDER CROWLEY METHOD WITH FLUX CORRECTION	WIND, DIFFUSIVITY, AND CONCENTRATION FIELDS	UPDATED POLLUTANT CONCENTRATION FIELD
CHEM2	CALCULATES THE $\text{SO}_2 \rightarrow \text{SO}_4^{2-}$ TRANSFORMATION VIA A FACILITATED DECAY MODEL	POLLUTANT CONCENTRATIONS, FLUXES, AND SOURCES	UPDATED POLLUTANT CONCENTRATIONS
CHEM5	SOLVES THE GRC PHOTOCHEMICAL MODEL USING 5 ACTIVE CHEMICAL SPECIES	POLLUTANT CONCENTRATIONS, FLUXES, AND SOURCES	UPDATED POLLUTANT CONCENTRATIONS
CHEM15	SOLVES THE REACTANT-SET-SEGMENT PHOTOCHEMICAL MODEL USING 14 ACTIVE SPECIES	POLLUTANT CONCENTRATIONS, FLUXES, AND SOURCES	UPDATED POLLUTANT CONCENTRATIONS
OUTPUT	PRODUCES A SUMMARY OF THE FIELD ARRAYS, AND/OR PRINTS SPECIES AS SPECIFIED IN THE INPUT OPTION DATA	FIELD ARRAYS, AND, DEPOSITION, STABILITY, CONCENTRATION	PRINTS AND CONTOURS OF FIELD ARRAYS
FINISH	ENDS THE PROGRAM	NONE	PRINTS END OF PROGRAM MESSAGE

2.1 COMMON

There are three blocks of common data used in IMPACT. The first block, /SCALER/, contains all the scalar quantities shared between subroutines. The second block, /ONETWO/, contains all the one and two-dimensional arrays that remain invariant from problem to problem. These common blocks are found in all subroutines except ADH, CHEM3, CHEM6, CHEM15 and FINISH. The third common stock, /TREFOR/, contains the two, three and four-dimensional arrays that may change size from problem to problem. Rather than include this common stock in every subroutine, the common appears only in the main program and is passed to other subroutines via the subroutine call statement. Thus, the user can vary the size of the problem with a minimum of effort. (See section 4.1 for a discussion on modifying the /TREFOR/ common block.)

A glossary of common variables is given in Table 2.

Table 2. Definition of Variables Used in COMMON.

COMMON/SCALAR/

DX = cell size, x-direction
DY = cell size, y-direction
DZ = cell size, z-direction
NX = number of cells, x-direction
NY = number of cells, y-direction
NZ = number of cells, z-direction
NXP1 = NX+1
NYP1 = NY+1
NZP1 = NZ+1
NXM1 = NX-1
NYM1 = NY-1
NZM1 = NZ-1
NUMEWD = number of wind stations
TIMEND = (not currently implemented)
NUMCEM = number of chemicals
NUMESD = number of stability profile locations
NUMHRS = number of hours simulated
NUMEPS = number of point sources
NUMASD = number of area sources
NOWTIM - current hour
IDOWND = wind model flag
IDOCEM = chemistry model flag
IDODIF = diffusion model flag
IDOPLT = surface plot flag
NUMBCP = number of background concentration points
TGRND = temperature of ground
SOLAR = flux
PRESR = ambient pressure
AMBCO = CO concentration, ppm
AMH2O = water vapor concentration, ppm
SIMTIM = same as NOWTIM
IDOPRN = print flag
IDOCAL = Calcomp plotter flag
IDT = number of time steps per hour
HRSAVG = number of hours to be averaged
DXI2 = $1/DX^2$
DYI2 = $1/DY^2$
DZI2 = $1/DZ^2$
DXI = $1/DX$
DYI = $1/DY$
DZI = $1/DZ$
NS = number of species
NUMSRD = number of surface roughness data
IDOPLM = plume rise model flag
DT = time step size
IDOSUR = surface plot flag

DX2 = DX^2

DY2 = DY^2

DZ2 = DZ^2

NOWEWD = time at which next wind station data read

NOWESD = time at which next stability profile data read

NOWEPS = time at which next point source data read

NOWASD = time at which next area source data read

NOWBCP = time at which next background concentration
data read

COMMON/ONETWO/

EPS = point source emission array

UDH = specified plume rise array

DH = plume rise array

CC = cellular concentration

SS = cellular source array

ISKIP = subroutine control flag array

IEW = I-location of wind data

JEW = J-location of wind data

EWS = wind speed array

EWD = wind direction array

EWW = wind-weight array

WV = measured wind speed, v-direction

IES = I-location of stability data

JES = J-location of stability data

ESS = stability profile array

WU = measured wind speed, u-direction

IBC = I-location of background concentration

JBC = J-location of background concentration

EBC = background concentration array

IPS = I-location of source

JPS = J-location of source

KPS = K-location of source

IZO = I-location of surface roughness

HPS = stack height array

TPS = stack gas temperature array

VPS = stack volume flow array

DPS = stack diameter array

SPS = exhaust velocity of stack

ISS = I-location of stack

JSS = J-locations of stack

SSS = (not implemented)

TITLE = title of run

TITL = titles of field arrays

JZO = J-location of surface roughness

ZO = surface roughness array

ILAS = low I-location of area source

IHAS = high I-location of area source

JLAS = low J-location of area source

JHAS = high J-location of area source

EAS = area source emission array

CEMNAM = chemistry array
EPST = point source emission array
EAST = area source emission array

COMMON /TREFOR/

C = concentration array
CA = averaged surface concentration array
CT = temporary concentration array
U = u-velocity
V = v-velocity
W = w (vertical) velocity
EX = horizontal diffusivity x-direction
EY = horizontal diffusivity y-direction
EZ = vertical diffusivity z-direction
STAB = stability array
HT = terrain height array
ZO = surface roughness array
IHT - integerized height array

2.2 WEST WIND FIELD MODEL - SUBROUTINE MARIAH

IMPACT is designed to be used with as little or much data as is available, since possible applications range from unmonitored wilderness to intensively instrumented research areas. Thus, the wind field uses any and all available data, with its accuracy improving with increased, usable data.

The wind field development is diagramed in Figure 3. Atmospheric stability data are interpolated via a $1/r^4$ weighting to produce a stability field over the computational domain (1.0 = A stability 7.0 = G stability). Wind observations are projected upwards from the point measured through the portion of the grid without any measurement. Such projected data are weighted as 0.001 compared with user-specified weighting of from 1.0 to 0.0.

The projections are based on a power law extrapolation of the form

$$S = S_0 (Z/Z_0)^P$$

for speeds S and S_0 at vertical coordinates Z and Z_0 , respectively, with P determined by linear interpolation from the following table:

Stability Category	A 1	B 2	C 3	D 4	E 5	F 6	G 7
P	.15	.17	.20	.26	.39	.48	.54

The maximum wind speed allowed is 20 m/sec and the initial Z_0 is assumed equal to $\Delta Z/2$. After the wind speeds are projected upwards, the wind speed and direction are resolved into U and V vector components at every wind data site. Next, the entire wind field is developed via a weighted $1/r^2$ interpolation of the vertical profiles for each horizontal plane. Up to this point, terrain has not been considered. Terrain first enters by shifting

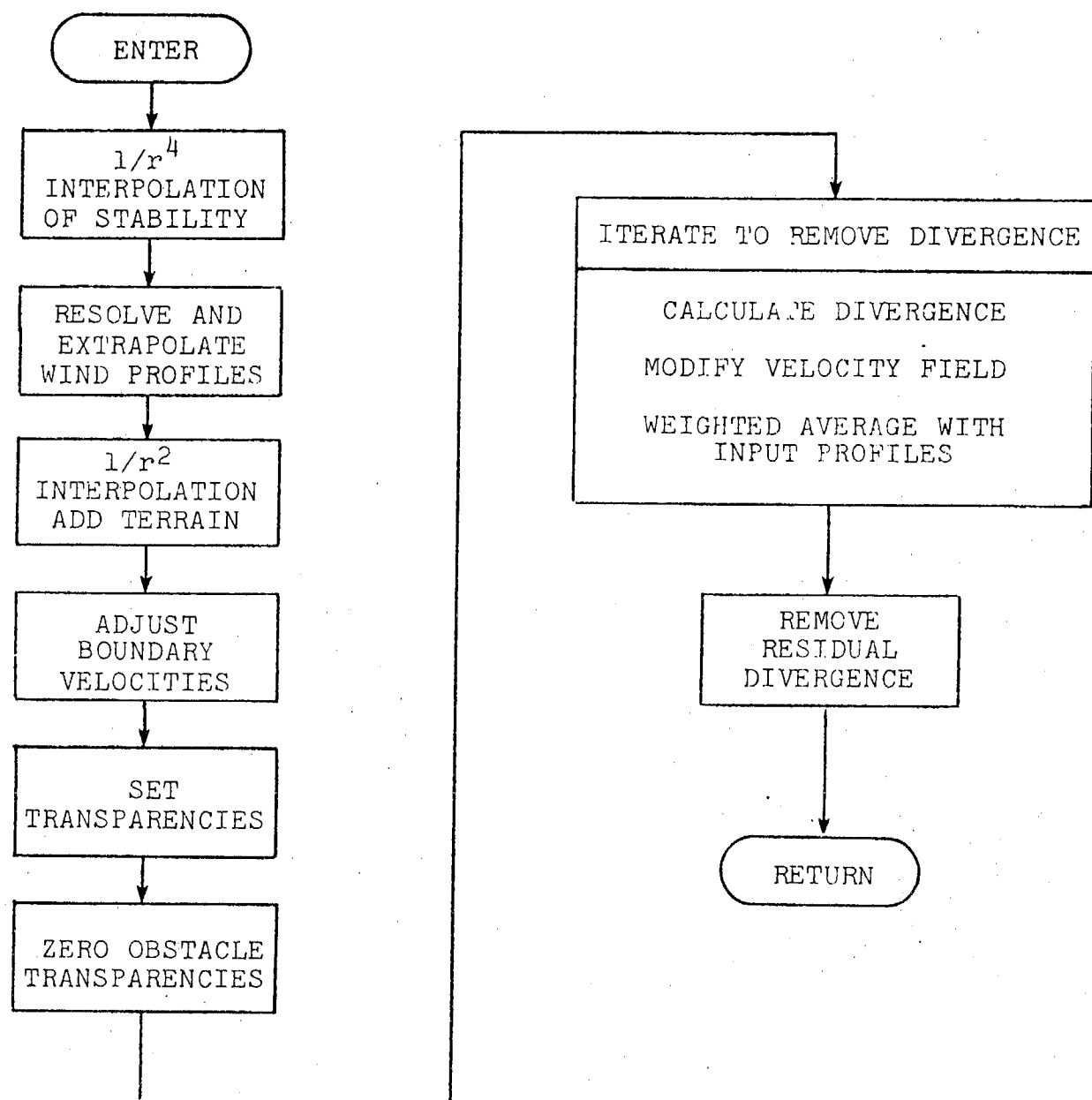


Figure 3. MARIAH Flow Diagram.

The subroutine uses eight corners for profiles and profiles are incorporated in the averaging of the iterative cycles, a feature of the grid. This

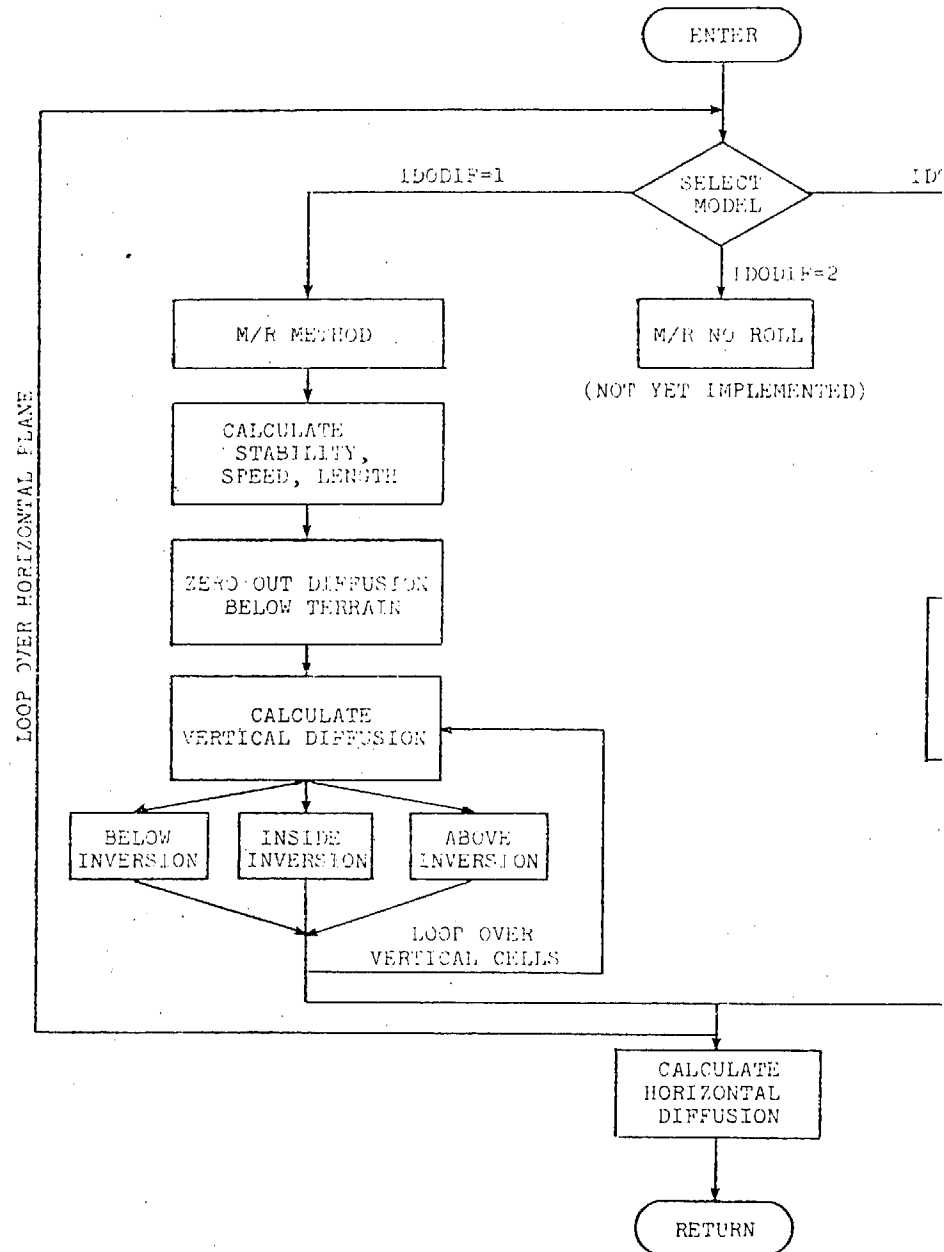


Figure 4. DIFFUS Flow Diagram.

where

Z is the height above the ground
 L is the Monin-Obukhov length and
 U_* is the friction velocity

$$U_* = (0.35) U_O / \{m(Z_w/Z_O) + 4.7 (Z_w - Z_O)/L\} \text{ for } L \geq 0$$

and

$$U_* = (0.35) U_O / \{m((Z'_w - 1.0)/(Z'_w + 1.0)) + 2.0 \tan^{-1}(Z'_w) - ((Z'_O - 1.0)/(Z'_O + 1.0)) - 2.0 \tan^{-1}(Z'_O)\} \text{ for } L < 0$$

$$Z'_w = (1.0 - 15 Z_w/L)^{1/4}$$

$$Z'_O = (1.0 - 15 Z_O/L)^{1/4}$$

U_O is wind speed at a height Z_w above the surface and
 $Z_w = \Delta Z/2$

The vertical diffusivity D is further modified by the function

$$D = D + (1.1 - Z/ZIBSE)$$

when

$$0.1 * ZIBSE < Z < 1.1 * ZIBSE$$

For the second case (inside an elevated inversion), the surface wind speed is set to 3.0 m/sec, the Monin-Obukhov length is set to 20.0, and Z (the height above the surface) is measured from ZIBSE.

Above the elevated inversion, the third case, neutral stability is assumed and Z is measured from the top of the inversion ZITOP. The vertical diffusion is then limited by $0.001 \text{ m}^2/\text{sec} \leq D \leq 1000 \text{ m}^2/\text{sec}$ and assigned to the top face of the cell.

The vertical diffusivity as calculated by the DEPICT model (Smith, 1971) is more straightforward.

$$D = (.45) \bar{U} \sigma_{\epsilon} l$$

where

\bar{U} is the wind speed at the point of interest
 σ_{ϵ} is the standard deviation of the wind vane fluctuation - σ_{ϵ} is empirically related to stability as follows:

Stability	1	2	3	4	5	6
σ_{ϵ}	.262	.237	.184	.119	.056	.023

l is the turbulence length scale (meters) and is empirically related to stability as follows:

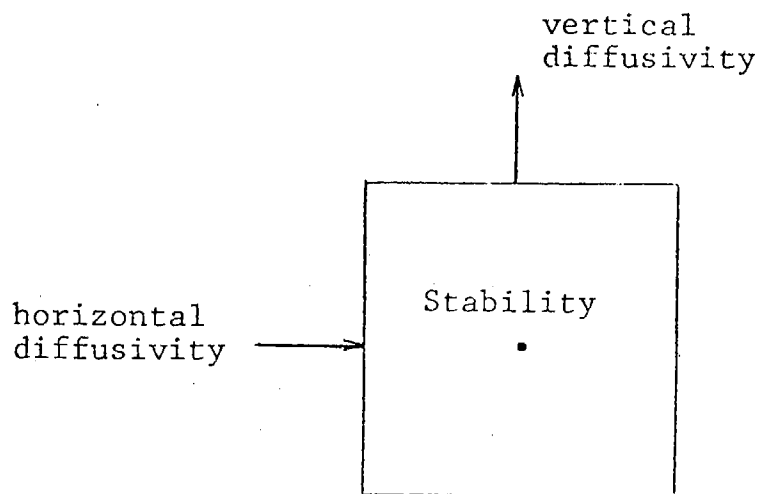
Stability	1	2	3	4	5	6
l	105	85	74	64	59	54

The horizontal diffusivities are calculated for either diffusion option using the relation $D_H = \alpha D_V$ where α depends on stability as follows:

Stability	1	2	3	4	5	6
α	.5	.75	1.05	1.35	1.70	2.0

(See Figure 4-11 in the main report.)

The location of the related stability vertical diffusivity and horizontal diffusivity parameters are indicated below:



Because the diffusivities are face-centered, the vertical stability is offset one-half cell above the cell center; the horizontal diffusivity is offset one-half cell to the left of the cell center.

2.4 PLUME RISE MODELS - SUBROUTINE PLUMER

The user is given the option of selecting the Briggs 1974 plume rise model, the Briggs 1967 plume rise model, or the TVA plume rise model; or specifying either $U(\Delta h)$ or Δh . A flow diagram of PLUMER is shown in Figure 5. The subroutine loops over each source and checks whether or not the source is inside the grid. If the source is outside the grid, an error message is printed and that source is ignored. If the user has specified ΔH (IDO PLM = 8), the total plume rise ($H + \Delta H$) is determined and the cell in which to insert the point source emissions is calculated. If $U(\Delta h)$ is specified (IDOPLM = 7), the velocity at the top of the stack is calculated and $\Delta H = U\Delta h/U$. For either Briggs model (IDOPLM = 1,2) and the TVA model (IDOPLM = 3), the buoyancy parameter F and the temperature gradient at the top of the stack are determined. The plume rise Δh is calculated as described in the following paragraphs.

TVA 1974

$$\text{Buoyancy } F = gV (T_o - T_a)/T_o$$

where

g is the gravitation acceleration (9.8 m/sec^2)

V is the initial volume flux divided by π (m^3/sec)

(also equal to $w r^2$ where w is the exit velocity
and r is the stack inside radius)

T_o is the stack gas temperature ($^{\circ}\text{K}$)

T_a is the ambient temperature at the stack ($^{\circ}\text{K}$) for
neutral and stable conditions.

For a distance of 1824 meters (a presumed final rise)
the plume rise is calculated from

$$\Delta h = 173 \frac{F^{1/3}}{u} \exp (-64 (\partial T / \partial z + 0.01))$$

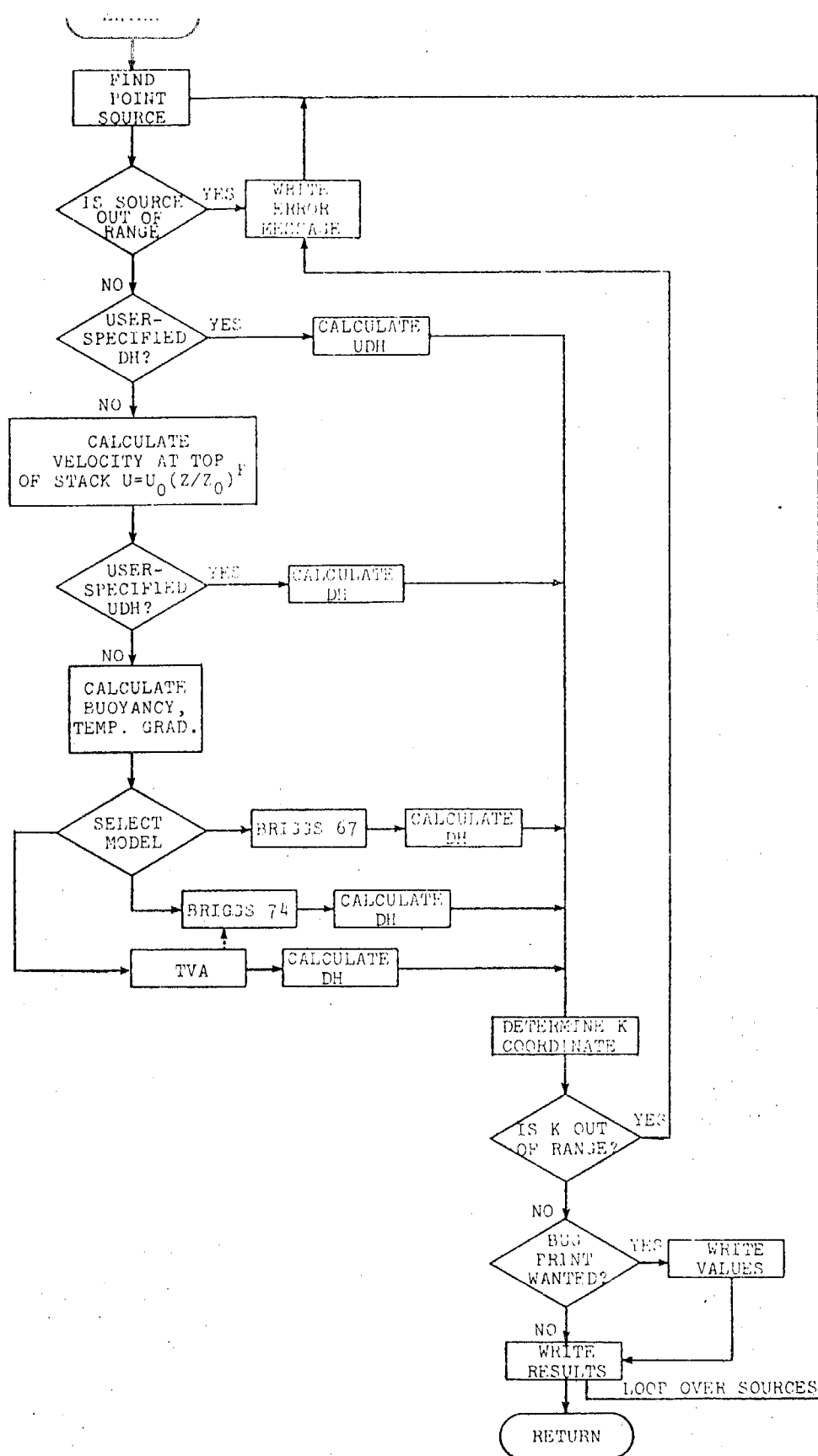


Figure 5. Plumer Flow Diagram.

Briggs 1967

Neutral and Unstable (A-D)

$$\Delta h = 1.6 \frac{F^{1/3}}{u} (10 h_s)^{2/3}$$

where h_s is the stack height

Stable Conditions (E-F)

$$\Delta h = 2.9 \left(\frac{F}{u s}\right)^{1/3} \text{ where } s = \frac{g}{T_a} \left(\frac{\partial T}{\partial z} + 0.01\right)$$

Briggs 1974

Neutral and Unstable (A-D)

$$\Delta h = 1.6 \frac{F^{1/3}}{u} (3.5 x^*)^{2/3}$$

where

$$x^* = 14 F^{5/8} \text{ for } F \leq 55$$

$$x^* = 34 F^{2/5} \text{ for } F > 55$$

Stable Conditions (E-F)

$$\Delta h = 2.6 \left(\frac{F}{u s}\right)^{1/3} \text{ where } s = \frac{g}{T_a} \left(\frac{\partial T}{\partial z} + 0.01\right)$$

2.5 POLLUTANT TRANSPORT - SUBROUTINES POLLUT, ADH

The transport, dispersion and transformation of pollutants is controlled by the subroutine POLLUT. Each call to POLLUT advances the simulation one hour. A flow diagram of POLLUT is shown in Figure 6.

The advection/diffusion time step is calculated such that the maximum normalized flux across any cell is less than alpha, i.e.:

$$\Delta t = \alpha / (u/\Delta x + kx/\Delta x^2)$$

If the time step is too large or too small, an error message is printed and the program stops. The time step is further adjusted such that N advection cycles end exactly on the hour.

The subroutine then loops over N time steps

$$(\Delta t \cdot N = 3600 \text{ seconds})$$

If multiple species chemistry is requested, POLLUT checks to see if each cell is either below terrain or contains a significant amount of tracer concentration. If not, the background concentration is loaded into the cell. This procedure forces those cells not affected by point or area sources to agree with the observed background concentration distribution.

The concentrations are stored in a temporary array and then advected/diffused in the X, Y, Z direction via successive calls to subroutine ADH. Point and area source emissions are added. If multiple species chemistry is selected, the updated concentrations in each cell are subtracted from the stored (old) concentrations to determine the flux into each cell. If there is sufficient tracer concentration in the cell ($> 10^{-3} \mu\text{g}/\text{m}^3$), the appropriate chemistry subroutine (CHEM 3, CHEM 6, CHEM 15) is called and returns with the updated concentration.

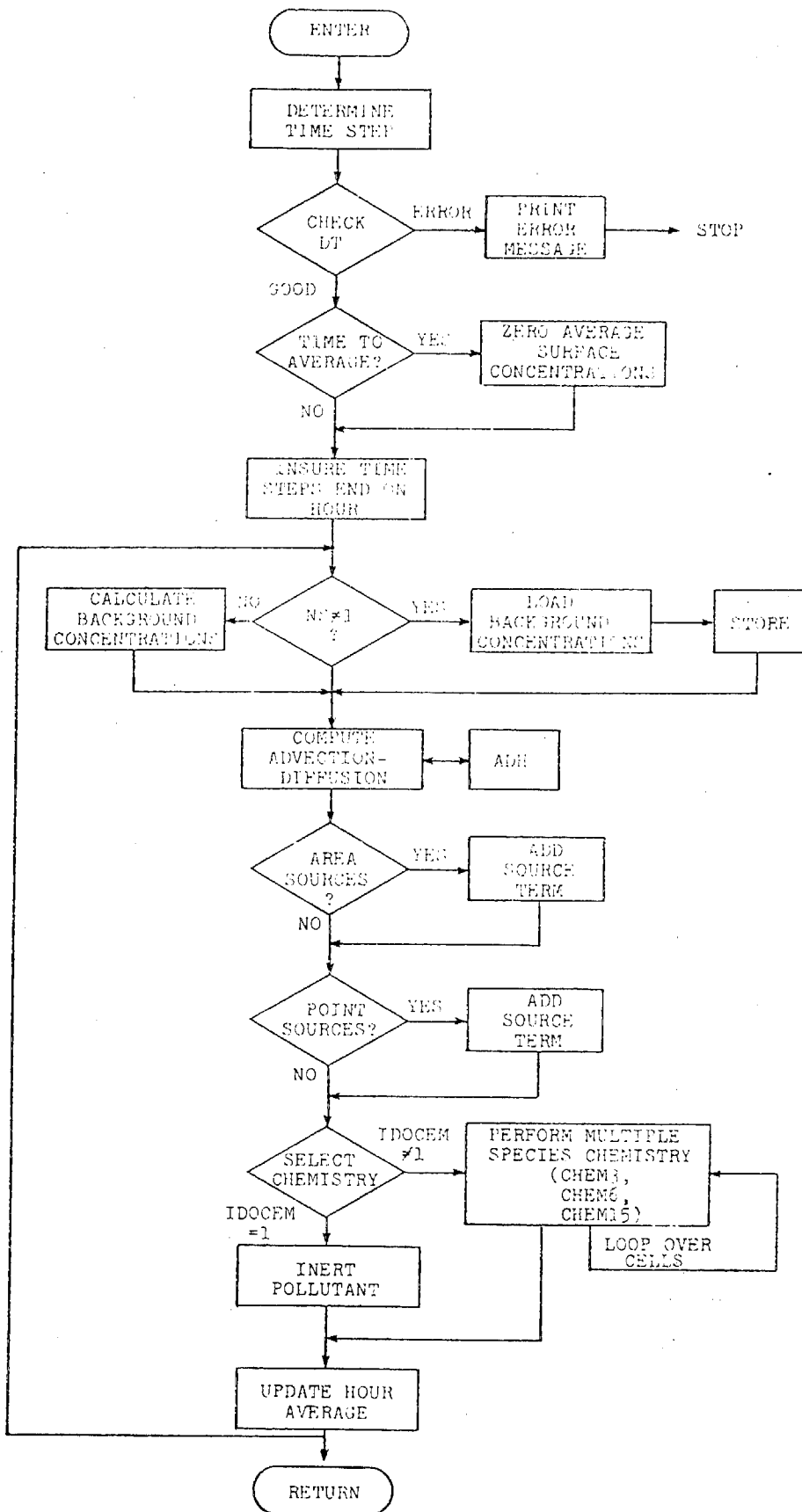


Figure 6. POLLUT Flow Diagram.

Finally, surface concentrations are accumulated to calculate the hour-averaged concentration field.

Subroutine ADH solves the differential equation describing advection and diffusion of pollutant concentration, i.e.:

$$\frac{\partial C}{\partial t} = -\nabla \cdot UC + \nabla \cdot K \cdot \nabla C$$

The solution is performed separately in each direction with the form

$$\frac{\partial C}{\partial t} = -\frac{\partial UC}{\partial x} + \frac{\partial}{\partial x} K_x \frac{\partial C}{\partial x}$$

The finite difference solution is based on a flux-corrected version of Crowely's second-order advection scheme

$$C_i^{n+1} = C_i^n + \frac{F_{i-\frac{1}{2}}^n - F_{i+\frac{1}{2}}^n}{\Delta x} \Delta t$$

where

$$F_{i+\frac{1}{2}}^n = \frac{\Delta x}{2\Delta t} \left\{ (C_i^n + C_{i+1}^n) \alpha_{i+\frac{1}{2}} + (C_i^n - C_{i+1}^n) \alpha_{i+\frac{1}{2}}^2 \right\} \\ + K_{xi+\frac{1}{2}} (C_i^n - C_{i+1}^n) / \Delta x$$

and

$$\alpha_{i+\frac{1}{2}} = \frac{U_{i+\frac{1}{2}} \Delta t}{\Delta x}$$

The flux corrector prohibits the calculation of negative concentrations by an adjustment of the flux, as follows:

if

$$F_{i+\frac{1}{2}} \geq 0$$

then

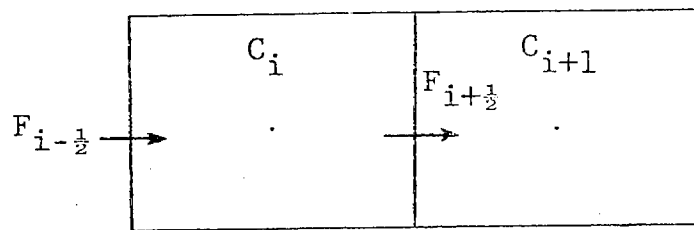
$$F_{i+\frac{1}{2}} = \min (F_{i+\frac{1}{2}}, \frac{\Delta x}{\Delta t} C_i, \frac{\Delta x}{\Delta t} C_i + F_{i-\frac{1}{2}})$$

if

$$F_{i+\frac{1}{2}} < 0$$

then

$$F_{i+\frac{1}{2}} = \min (-F_{i+\frac{1}{2}}, \frac{\Delta x}{\Delta t} C_{i+1})$$



2.6 CHEMISTRY - SUBROUTINES CHEM 3, CHEM 6, CHEM 15

The IMPACT code is designed to provide the framework for simulating the complex chemical transformations of air pollutants, as well as the transport and dispersion of simple inert pollutants. Three chemical reaction mechanisms are available as options in the present version of IMPACT. The input format to all chemistries are the same: the cellular concentrations before the current advection/diffusion step, $C(N)$; the input (or output) flux due to the current advection, diffusion, and emissions, $\epsilon(N)$; and the advection/diffusion time step, Δt . In addition, the two photochemistry models require the solar flux, SOL, as input. All chemistries return the pollutant concentrations at the new time $T = T + \Delta t$ via the cellular concentration array $C(N)$.

Two chemical reaction mechanisms simulate the transformation of the constituents of photochemical smog. A simple mechanism developed by GRC (Eschenroder, 1972), CHEM6 supports five active species including NO_2 , NO , O_3 , and HC (non-methane reactive hydrocarbon and HN_2O_2). CHEM6 is limited to simulations of 12 hours or less (see Table 3). The other photochemical model, CHEM15, incorporates the latest EPA-recommended mechanisms (Hect-Seinfeld, 1974), and supports 14 active species including NO_2 , NO , O_3 , HCl (olefins), $HC2$ (aromatics), $HC3$ (parafins), $HC4$ (aldehydes), N_2O_5 , HN_2O_3 , HN_2O_2 , H_2O_2 , R_00 , and $RCOO$ (see Table 4).

Both CHEM6 and CHEM15 express the effects of chemical reactions and sources by a solution for each cell of a rate equation of the form

$$\frac{\partial C_1}{\partial t} = A_1 + \sum_{n=1}^N B_{1n} C_n + \sum_{n,m=1}^N D_{1nm} C_n C_m$$

Table 3. CHEM6 Reaction Mechanism.

	<u>REACTION MECHANISM</u>	<u>RATE CONSTANT (ppm/min)</u>
1.	$h\nu + NO_2 \rightarrow NO + O$	2.67×10^{-1}
1a.	$O + O_2 + M \rightarrow O_3 + M$	$2. \times 10^6$
2.	$NO + O_3 \rightarrow NO_2 + O_2$	2.67×10
3.	$O + HC \rightarrow (b_1) RO_2$	2.67×10^2
4.	$OH + HC \rightarrow (b_2) RO_2$	$1. \times 10^4$
5.	$O_3 + HC \rightarrow (b_3) RO_2$	$4. \times 10^{-3}$
6.	$RO_2 + NO \rightarrow NO_2 + (y) OH$	$1. \times 10^5$
7.	$RO_2 + NO_2 \rightarrow PAN$	$2. \times 10^2$
8.	$OH + NO \rightarrow HONO$	1.5×10^3
9.	$OH + NO_2 \rightarrow HNO_3$	$3. \times 10^3$
10.	$h\nu + HONO \rightarrow OH + NO$	1.5×10^{-3}
	H_2O	
11.	$NO + NO_2 \rightarrow 2 HONO$	$1. \times 10^{-3}$
12.	$NO_2 + O_3 \rightarrow NO_3 + O_2$	$5. \times 10^{-3}$
13.	$NO_3 + NO_2 \rightarrow N_2O_5$	4.5×10^3
14.	$N_2O_5 \rightarrow NO_3 + NO_2$	1.4×10
15.	$N_2O_5 + H_2O \rightarrow 2HNO_3$	6.05×10
16.	$NO_2 + PARTICULATES \rightarrow PRODUCTS$	0.0

Table 4. CHEM15 Reaction Mechanism.

REACTION MECHANISM	RATE CONSTANT (ppm/min)
1. $\text{NO}_2 + h\nu \rightarrow \text{NO} + \text{O}$	2.66×10^{-1}
2. $\text{O} + \text{O}_2 + \text{M} \rightarrow \text{O}_3 + \text{M}$	$2. \times 10^{-5}$
3. $\text{O}_3 + \text{NO} \rightarrow \text{NO}_2 + \text{O}_2$	2.08×10^1
4. $\text{O} + \text{NO} + \text{M} \rightarrow \text{NO}_2 + \text{M}$	3.5×10^{-3}
5. $\text{O} + \text{NO}_2 \rightarrow \text{NO} + \text{O}_2$	1.38×10^4
6. $\text{O} + \text{NO}_2 + \text{M} \rightarrow \text{M} \rightarrow \text{NO}_3 + \text{M}$	2.2×10^{-3}
7. $\text{O}_3 + \text{NO}_2 \rightarrow \text{NO}_3 + \text{O}_2$	4.6×10^{-2}
8. $\text{NO}_3 + \text{NO} \rightarrow 2\text{NO}_2$	1.5×10^4
9. $\text{NO}_3 + \text{NO}_2 \rightarrow \text{N}_2\text{O}_5$	4.5×10^3
10. $\text{N}_2\text{O}_5 \rightarrow \text{NO}_2 + \text{NO}_3$	1.5×10
11. $\text{N}_2\text{O}_5 + \text{H}_2\text{O} \rightarrow 2\text{HNO}_3$	$1. \times 10^{-5}$
12. $\text{NO} + \text{HNO}_3 \rightarrow \text{HNO}_2 + \text{NO}_2$	2.5×10^{-4}
13. $\text{HNO}_2 + \text{HNO}_3 \rightarrow \text{H}_2\text{O} + 2\text{NO}_2$	$2. \times 10^{-1}$
14. $\text{NO} + \text{NO}_2 + \text{H}_2\text{O} \rightarrow 2\text{HNO}_2$	2.1×10^{-6}
15. $2\text{HNO}_2 \rightarrow \text{NO} + \text{NO}_2 + \text{H}_2\text{O}$	4.5
16. $\text{HNO}_2 + h\nu \rightarrow \text{OH} + \text{NO}$	1.3×10^{-2}
17. $\text{OH} + \text{NO}_2 \rightarrow \text{HNO}_3$	1.5×10^4
18. $\text{OH} + \text{NO} + \text{M} \rightarrow \text{HNO}_2 + \text{M}$	1.2×10^4
19. $\text{OH} + \text{CO} + (\text{O}_2) \rightarrow \text{CO}_2 + \text{HO}_2$	2.5×10^2
20. $\text{HO}_2 + \text{NO} \rightarrow \text{OH} + \text{NO}_2$	$7. \times 10^2$
21. $\text{H}_2\text{O}_2 + h\nu \rightarrow 2\text{OH}$	2.5×10^2
22. $\text{HC}_1 + \text{O} \rightarrow \text{ROO} + \alpha\text{RCOOO} + (1-\alpha) \text{HO}_2$	6.8×10^3
23. $\text{HC}_1 + \text{O}_3 \rightarrow \text{RCOOO} + \text{RO} + \text{HC}_4$	1.6×10^{-2}
24. $\text{HC}_1 + \text{OH} \rightarrow \text{ROO} + \text{HC}_4$	2.5×10^4
25. $\text{HC}_2 + \text{OH} \rightarrow \text{ROO} + \text{OH}$	1.07×10^2
26. $\text{HC}_2 + \text{OH} \rightarrow \text{ROO} + \text{H}_2\text{O}$	$8. \times 10^3$
27. $\text{HC}_3 + \text{O} \rightarrow \text{ROO} + \text{OH}$	6.5×10
28. $\text{HC}_3 + \text{OH} \rightarrow \text{ROO} + \text{H}_2\text{O}$	3.8×10^4
29. $\text{HC}_4 + h\nu \rightarrow \beta\text{ROO} + (2-\beta) \text{HO}_2$	2.5×10^{-3}
30. $\text{HC}_4 + \text{OH} \rightarrow \beta\text{RCOOO} + (1-\beta) \text{HO}_2 + \text{H}_2\text{O}$	2.3×10^4
31. $\text{ROO} + \text{NO} \rightarrow \text{RO} + \text{NO}_2$	9.1×10^2
32. $\text{RCOOO} + \text{NO} + (\text{O}_2) \rightarrow \text{ROO} + \text{NO}_2 + \text{CO}_2$	9.1×10^2
33. $\text{RCOOO} + \text{NO}_2 \rightarrow \text{RCOOO NO}_2$	$1. \times 10^2$
34. $\text{RO} + \text{O}_2 \rightarrow \text{HO}_2 + \text{HC}_4$	2.4×10^{-2}
35. $\text{RO} + \text{NO}_2 \rightarrow \text{RONO}_2$	4.9×10^2
36. $\text{RO} + \text{NO} \rightarrow \text{RONO}$	2.5×10^2
37. $\text{HO}_2 + \text{HO}_2 \rightarrow \text{H}_2\text{O}_2 + \text{O}_2$	5.3×10^3
38. $\text{HO}_2 + \text{ROO} \rightarrow \text{RO} + \text{OH} + \text{O}_2$	$1. \times 10^2$
39. $2\text{ROO} \rightarrow 2\text{RO} + \text{O}_2$	$1. \times 10^2$

HC_1 = olefins, HC_2 = aromatics, HC_3 = paraffins, HC_4 = aldehydes

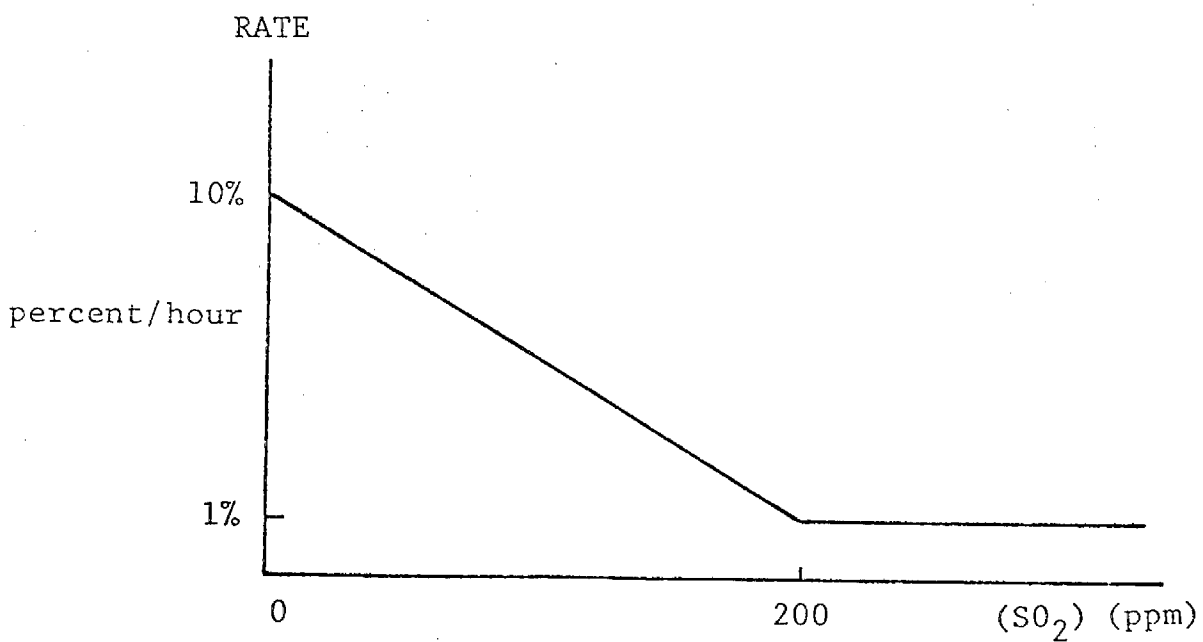
for the 1th pollutant species. The A_1 factor is used to enter sources and the net advective/diffusive flux into the cell. The rate equation is recast into

$$\frac{\partial C_1}{\partial t} = F_1 - R_1 C_1$$

and an approximate solution, used as a predictor-corrector, is

$$C_1^{n+1} = - \frac{F_1}{R_1} + \left(C_1^n + \frac{F_1}{R_1} \right)^{-R_1 \Delta t}$$

The third chemistry option (CHEM 3) is a simplified treatment of the oxidation of sulfer dioxide to sulfate. The SO_2 mechanism currently encoded in IMPACT does not simulate SO_2 photochemistry but rather relies upon empirical observation of atmospheric SO_2 oxidation rates. Observed maximum rates have been found to range from 2 percent per hour to 12 percent per minute. The SO_2 mechanism in IMPACT employs a spatially-dependent SO_2 to SO_4 conversion rate that depneds upon the plume concentration at the point under consideration as shown below.



2.7 INPUT/OUTPUT ROUTINES

Four subroutines control the input and output of IMPACT. Subroutine READEM reads and writes the initial invariant input data. Subroutine SUMMARY reads and processes the terrain and surface roughness data and prints a summary of the invariant data. Subroutine METSOR reads the hourly wind, stability, point source, area source, background concentration, and global parameters (solar flux, surface temperature, etc.). Emissions data (except for the tracer) are transformed from g/sec to ppm/sec and a summary of the hourly input data is printed. Subroutine OUTPUT controls the printing of the results of the IMPACT calculations. Printer edits and contours of winds, diffusivity, stability, and concentration arrays can be printed level-by-level or for surface values only. The surface array of hour-average concentrations can also be printed.

Subroutine CONTUR creates a printer contour plot from any submitted array with less than 45 cells in the x-direction. Subroutine PRINIT prints the first two significant digits of any submitted array with less than 40 cells in the x-direction.

3. PROGRAM OPTIONS

IMPACT allows the user to select the size and resolution of the simulation problem. However, due to the size of dimensioned arrays in the COMMON/ONETWO/ block and the formats in the output subroutines CONTUR and PRINIT, the following limitations apply:

- NX - the number of cells in the x-direction must be ≤ 40
- NZ - the number of cells in the y-direction must be ≤ 20
- NUMEWD, NUMESD, NUMEPS, NUMASD, NUMBCP and NUMSRD - the number of wind, stability, point source, area source, background, and surface roughness data sites (respectively) cannot be greater than 50 each.

Other than these conditions, only computer storage and run time considerations restrict the size and resolution of the problem. The time-dependent options allow the user to select the number of hours to be simulated (NUMHRS), and the interval (in multiples of one hour) for which time-averaged surface concentrations are calculated (HRSAVG). The user may also specify the time interval (in multiples of one hour) for the printer edit and printer contour output (IDOPRN, IDOPLT).

By selecting the appropriate input options, the user can specify the models to be used in calculating variables listed below:

IDOCM = 0	wind and diffusion field calculations only
IDOCM = 1	tracer only
IDOCM = 3	tracer plus SO_2 , SO_4 sulfate chemistry
IDOCM = 6	tracer plus photochemistry, GRC photochemical model
IDOCM = 15	tracer plus photochemistry, Hect-Seinfeld photochemical model
IDODIF = 1	Myrup/Ranzieri diffusivity model
IDODIF = 3	DEPICT diffusivity model

IDOPLM = 1 Briggs 1967 plume rise model
IDOPLM = 2 Briggs 1974 plume rise model
IDOPLM = 3 TVA plume rise model
IDOPLM = 7 user-specified $U\Delta h$ plume rise parameter
IDOPLM = 8 user-specified plume rise Δh .

4.2 INPUT FILE STRUCTURE

The input file for IMPACT is illustrated in Figure 7. Title, grid size, options, output, terrain, and surface roughness data are temporally invariant and are entered at the start of the simulation. Other inputs (wind, stability, point source, area source, background concentrations, and global atmospheric parameters) must be read in for each hour of simulation.

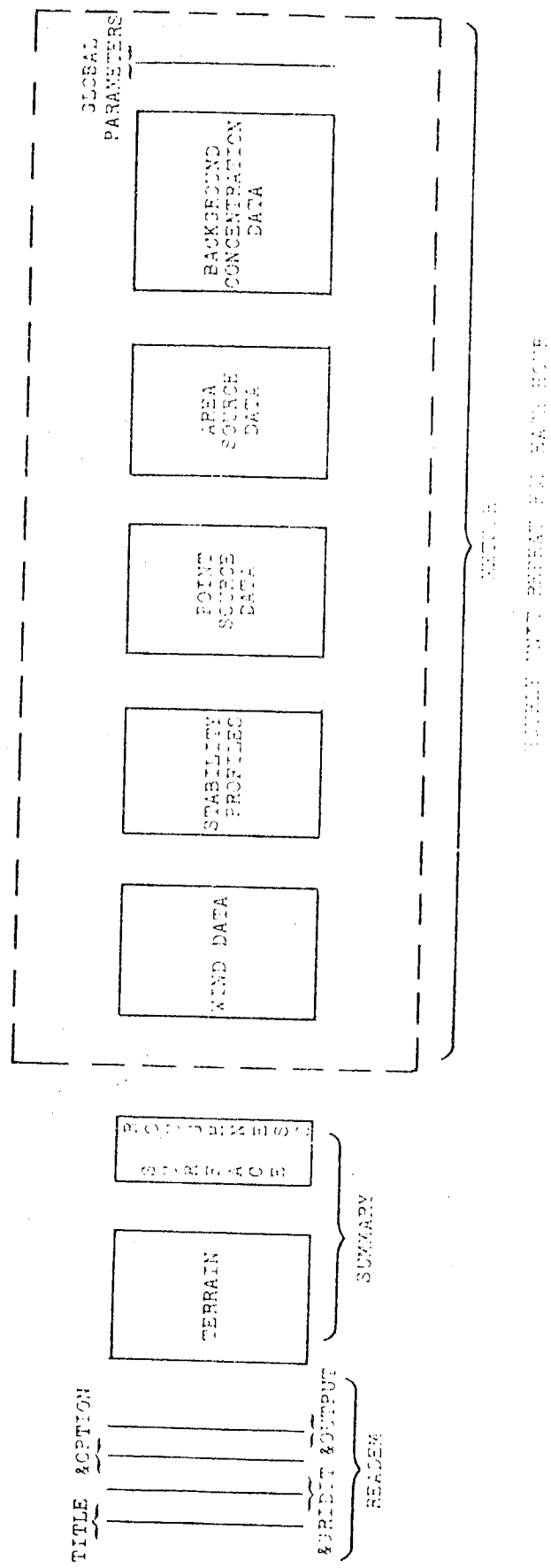


Figure 7. Data File Structure Indicating Calling Routines.

4.3 INPUT FILE DESCRIPTION

This section contains a record-by-record description of the input file required to run IMPACT.

Three records are read in using the NAMELIST convention (GRIDIT, OPTION, OUTPUT). The remaining records are transferred under format control.

The input records are described below in the order shown in Figure 7.

4.3.1 Title

The title record consists of up to 80 characters that identify the particular run read in under a 20A4 format.

4.3.2 & GRIDIT

This namelist record describes the size and resolution of the problem.

DX: } specifies the cell size in meters (east-west,
DY: } north-south, and vertical).
DZ: }
NX: } specifies the number of grid cells in the
NY: } x, y, z-directions.
NZ: }

4.3.3 & OPTION

This namelist record describes the options requested for wind field, chemistry, point source, plume rise and diffusion model

IDOWND: selects wind field model IDOWND = 1 WEST
IDOCEM: selects the chemistry model
 IDOCEM = 0 no pollutants, wind and diffusion field calculation only
 IDOCEM = 1 no chemistry, single tracer pollutant (NS = 1)

IDOCEM = 3 sulfate chemistry (NS = 3)
IDOCEM = 6 GRC photochemistry (NS = 6)
IDOCEM = 15 EPA photochemistry (NS = 15)
IDOPLM: selects the plume rise option
IDOPLM = 1 Briggs 69 plume rise
IDOPLM = 2 Briggs 74 plume rise
IDOPLM = 3 TVA plume rise
IDOPLM = 7 user-specified ΔH
IDOPLM = 8 user-specified ΔH
IDODIF: selects the diffusivity model option
IDODIF = 1 Myrup/Ranzieri diffusivity model
IDODIF = 3 DEPICT diffusivity model

4.3.4 & OUTPUT

This namelist record describes the total number of hours to be simulated and the output options desired.

NUMHRS: the number of hours to be simulated by IMPACT
IDOPLT: the frequency in integer hours for printer contour output
 IDOPLT = 0 no printer contour desired
 IDOPLT = 1 printer contours every hour
 IDOPLT = 3 printer contours every 3 hours plus first hour (i.e. hours 1, 3, 6...)
IDOPRN: the frequency in integer hours for printer edit output
 IDOPRN = 0 no printer edits desired
 IDOPRN = 1 printer edit every hour
 IDOPRN = 3 printer edit every 3 hours plus first hour
IDOCAL: option for calcomp plots of wind field vectors and pollutant concentration contours - not currently implemented, set IDOCAL = 0
HRSAVG: the integer number of hours desired for calculating average surface concentration
 HRSAVG = 1 one-hour average concentrations desired, printed every hour (IDOPRN, IDOPLT should also be set to 1)

HRSAVG = 3 three-hour average concentrations desired, printed every third hour (3, 6, 9...) IDOPRN, IDOPLT should be set to 1 or 3)

IDOSUR: option for prints all vertical levels of the field arrays or just the values at the surface

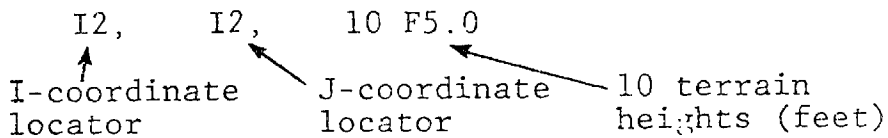
IDOSUR = 0 print all vertical levels plus the surface

IDOSUR = 1 print surface values only

4.3.5 Terrain Deck

Each card in the terrain deck holds consecutive x terrain heights (in feet) and is self-identified by having x and y cell location values. If an entire record contains zero heights for each location, it is not required. A record with a value of I = -1 is required to terminate the terrain deck.

Each terrain record holds up to 10 distinct elevations. The terrain record format is diagrammed below:



The I-coordinate locator value is the I-coordinate of the 1st terrain value on the record and can have only the following values: 1, 11, 21, 31. The values on the card thus correspond to locations at (I, J), (I+2, J), ... (I+9, J). The J-coordinate locator has the values 1, 2, 3... NY.

4.3.6 Surface Roughness Deck

The surface roughness deck contains two types of records. The first record, read as an I5 format, describes the number of surface roughness data records to follow. This information is stored in NUMSRD. The next NUMSRD records contain data on the location and value of surface roughness. The format is as follows:

I5, I5, F5.0
I-coordinate J-coordinate Surface roughness
(meters)

This data is only required if the Myrup/Ranzieri diffusivity model is desired. If no surface roughness data is needed, set NUMSRD = 0 to terminate the surface roughness deck.

The following data decks are required each hour except as noted.

4.3.7 Wind Data Deck

The first record of the wind data deck describes the number of wind data sites (stored as NUMEWD) and the hour at which the next wind data deck is to be read (stored as NOWEWD). This procedure allows the user to use the same wind data for multiple hours without rereading the same data deck each hour. If NOWEWD is set to 0, the wind data deck must be read in the next hour. If (at hour 2) NUMEWD = 5, the data deck read in at hour two will be used to calculate the wind field for hours 2, 3, and 4; at hour 5, a new wind data deck must be entered. The format of the data deck specification record is as follows:

I5, I5
number of wind data locations, NUMEWD hour at which next wind data deck is to be read, NOWEWD

For each wind data site, at least two records are required. The first record specifies the location of the wind data site and the confidence value (from 0.0 - 1.0) that the user places in the data (a value of 1.0 will force the wind field to agree with input value at the site, while a value of 0.0 will initialize the wind field using the input data but will allow the wind field to adjust itself without further regard to the input data). Valid, reliable hour average wind data is usually weighted as 1.0, while instantaneous pibal data is usually weighted as 0.1.

Note: The number of terrain cells at any location I, J can be determined by:

$$IHT(I,J) = \text{Integer portion of } \left[\frac{(HT(I,J) - HTMIN)}{3.2808 \cdot DZ} + .5 \right]$$

where

IHT (I,J) is the number of terrain cells at I,J

HT (I,J) is the height in feet at I,J

HTMIN is the minimum value in the HT array

DZ' is the vertical cell size in meters

At least one wind data location is required and a maximum of 50 data locations are permitted.

4.3.8 Stability Data Deck

The first record of the stability data deck, like the first record of the wind data deck, describes the number of stability data sites (stored as NUMESD) and the hour at which the next set of stability data is to be read (stored as NOWESD). The format is the same as for the wind data record:

I5, I5
number of stability data locations, NUMESD hour at which the next stability data is to be read, NOWESD

For each stability data location, at least two records are required. the first record contains the location of the stability data:

I5, I5
I-location J-location

The next record contains up to 16 values of atmospheric stability. If NZ > 16, another record containing stability data for K = 17, 18, 19, 20 is needed. The stability data is entered in the following format:

16 F5.0

where values of stability may be specified as:

- 0 inside terrain
- 1.0 - 1.5 Pasquill stability class A
- 1.5 - 2.5 Pasquill stability class B
- 2.5 - 3.5 Pasquill stability class C
- 3.5 - 4.5 Pasquill stability class D
- 4.5 - 5.5 Pasquill stability class E
- 5.5 - 6.5 Pasquill stability class F
- 6.5 - 7.0 Pasquill stability class G

Table 5 may be used as a guide for converting temperature sounding data to stability. Unlike the wind data, the stability data is not entered relative to local ground-level. The code does not fill in blanks in the vertical; thus, the user must enter exactly NZ values of stability (assigning zero to cells inside terrain and assigning stabilities to cells which actually may not have had a datum value).

Table 5. Classification of Atmospheric Stability
According to AEC Safety Guide 23.

Stability Classification	Pasquill Categories	Temperature Change With Height ($^{\circ}\text{C}/100\text{m}$)
Extremely unstable	A	< -1.9
Moderately unstable	B	-1.9 to -1.7
Slightly unstable	C	-1.7 to -1.5
Neutral	D	-1.5 to -0.5
Slightly stable	E	-0.5 to 1.5
Moderately stable	F	1.5 to 4.0
Extremely stable	G	> 4.0

Example: NZ = 17

K=	17	
	16	
	B	
	14	B
	13	
	12	F
	11	F
	10	
	9	E
	8	D
	7	
	6	
	5	
	4	
	3	
	2	
terrain	1	

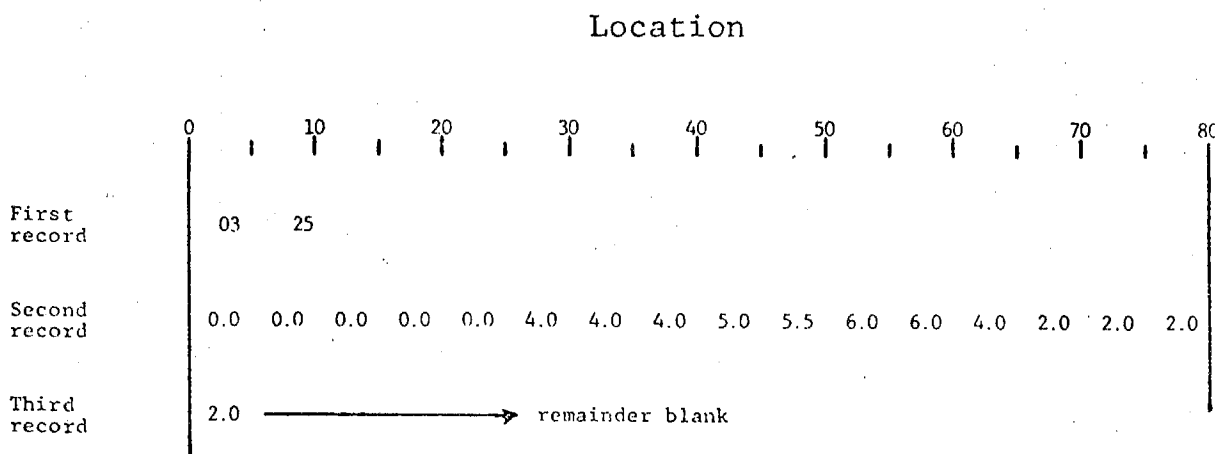
Location I = 3, J = 25

Set B = 2.0, F = 6.0, E = 5.0, D = 4.0

Extra values for needed cells 17, 16, 13, 10,
7, 6

Choose \longrightarrow 2.0, 2.0, 4.0, 5.5, 4.0,
4.0 respectively

The input records for this site would be:



At least one stability datum must be specified for every K-level; a maximum of 50 stability data sites are allowed.

4.3.9 Point Source Deck

If IDOCM = 0, the point source data deck is not required. The first record of the point source deck, like the first record of the wind data deck, describes the number of point sources (stored as NUMEPS), and the hour at which the next point source data is to be read (stored as NOWEPS). If only one set of point source data is desired, set NOWEPS to a number greater than the number of hours to be simulated (NUMHRS). The format is the same as the wind data record, i.e.,

I5, I5
↑ ↑
NUMEPS NOWEPS

If no point sources are needed for a particular hour, set NUMEPS = 0 and omit the following records.

For each point source, two records are required. The first record contains information on the stack and its operating conditions. The format and the associated variables are as follows:

I5, I5, F5.0, F5.0, F5.0, F5.0, F5.0, F5.0
↑ ↑ ↑ ↑ ↑ ↑ ↑ ↑
IPS JPS HPS TPS VPS DPS SPS UDH

Stack variable definitions

IPS: The I-coordinate of the stack

JPS: The J-coordinate of the stack

HPS: Stack height in meters (relative to local terrain)

TPS: Temperature of the plume in $^{\circ}\text{C}$

VPS: The stack volume flow in m^3/sec

DPS: The inside diameter at the top of the stack in meters

SPS: The exit plume velocity in meters/sec

UDH: A specified plume rise parameter

If

$$\text{IDOPLM} = 7 \quad \Delta h = \text{UDH/S} \quad \text{UDH in } m^2/\text{sec}$$

If

$$\text{IDOPLM} = 8 \quad \Delta h = \text{UDH} \quad \text{UDH in meters}$$

where

Δh is the plume rise (meters)

S is the wind speed at the top of the stack m/sec

If VPS is specified, DPS and SPS are not required, or vice versa.

If UDH is specified, then TPS, VPS, DPS and SPS are not required.

The second record contains information on the point source emissions. The input on this record varies with the choice of IDOCEM. However, all data are read into EPST in a 16F5.0 format and are in units of grams/second. They are specified as follows:

	<u>IDOCEM = 1</u>	<u>IDOCEM = 3</u>	<u>IDOCEM = 6</u>	<u>IDOCEM = 15</u>
EPST (1)	tracer	tracer	tracer	tracer
EPST (2)		SO_2	NO_2	NO_2
EPST (3)		SO_4	NO	NO
EPST (4)			O_3	O_3
EPST (5)			HC (reactive)	HC1 (olefins)
EPST (6)			HNO_2	HC2 (aromatics)
EPST (7)				HC3 (parafins)
EPST (8)				HC4 (aldehydes)
EPST (9)				N_2O_5
EPST (10)				HNO_3
EPST (11)				HNO_2
EPST (12)				HO_2
EPST (13)				H_2O_2
EPST (14)				ROO
EPST (15)				RCOO

A maximum of 50 point sources may be specified.

4.3.10 Area Source Deck

If IDOCM = 0, area source deck is not required.

The first record of the area source deck describes the number of area sources (stored as NUMASD) and the hour at which the next area source deck is to be read (stored as NOWASD). The format is the same as for the point source data:

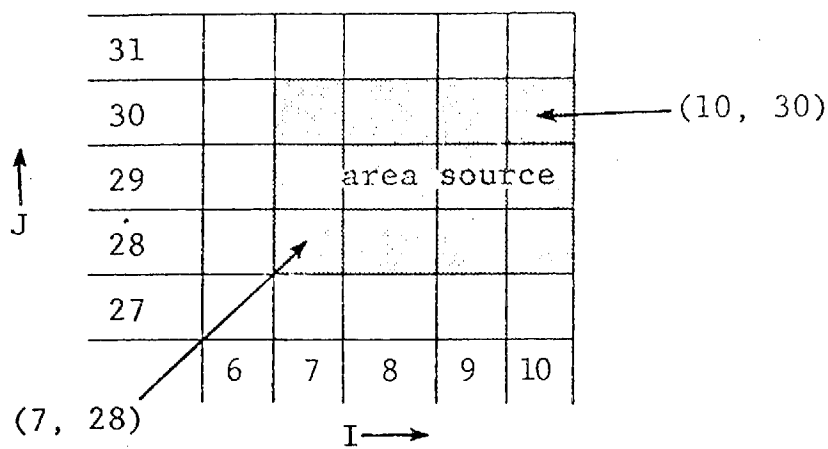
15, 15
↑ ↑
NUMASD NOWASD

If no area sources are needed for a particular hour, set NUMASD = 0 and omit the following records. If only one set of area source data is desired for the entire problem set NOWASD > NUMHRS.

For each area source, two records are required. The first record contains information on the location of the area source. Each area source is defined as a rectangle whose sides are an integral number of DX and DY. The locations of the lower-left corner and the upper-right corner of the area source are required to specify the area source. The format for this record is as follows:

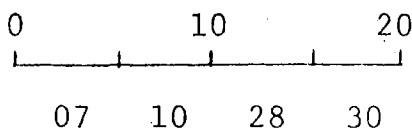
ILAS IHAS JLAS JHAS

Example:



In this example, ILAS = 7, IHAS = 10, JLAS = 28, JHAS = 30.

The record would look like this:



If only one cell is required to specify an area source, set ILAS = IHAS, JLAS = JHAS. However, all data must be entered in this record. The second record contains information on the area source emissions. The format of this record is exactly the same as the point source emissions record. (Refer to section 4.13.13 for further information.)

The emissions required for the area source deck are in grams per second per grid cell. Thus, if the emissions for the entire area source is E g/sec, the emissions per grid cell, ϵ g/sec/grid cell, equal $E / ((IHAS-ILAS + 1) \cdot (JHAS-JLAS + 1))$. The variable ϵ is required as input in this record.

A maximum of 50 area sources may be specified.

4.3.11 Background Concentration Deck

The background concentration deck allows the user to specify the spatial distribution of pollutants. Those cells not affected by point or area sources (defined as cells whose tracer concentrations is less than $0.001 \mu\text{g}/\text{m}^3$) are assigned concentrations based on a weighted, $1/r^2$, interpolation of background concentration values at the particular vertical level. Thus, the user must specify the vertical profile for all pollutants at each site. If IDOCM = 0, the background concentration deck is not required.

The first record of the background concentration deck describes the number of background sites (stored as NUMBCP) and the hour at which the next background concentration deck is to be read (stored as NOWBCP). The format is the same for the point source data:

I5, I5
NUMBCP NOWBCP

If no background data is needed, set NUMBCP and omit the following records. If only one set of data is needed for the entire problem, set NOWBCP > NUMHRS. If, for example, no data is needed for the entire problem (e.g., if IDOCDEM = 1, tracer only), set NUMBCP = 0 and NOWBCP = NUMHRS + 1.

For each background concentration data point, NZ + 1 records are required. The first record describes the coordinate location:

I5, I5
IBC JBC

where

IBC is the I-location of the background data

JBC is the J-location of the background data

The next NZ records describe the background concentrations of the reactive pollutants for each level starting at K = 1 and continuing to K = NZ. If, for example, levels K = 1 to K = 6 are below the surface, six blank cards must be inserted to account for levels K = 1 to K = 6. Every level must be specified. The input on these records varies with the selection of IDOCDEM. However, all data are read into EBC in a 14F5.0 format and are in units of ppm. They are specified as follows:

	<u>IDOCDEM = 3</u>	<u>IDOCDEM = 6</u>	<u>IDOCDEM = 15</u>
EBC (1)	SO ₂	NO ₂	NO ₂
EBC (2)	SO ₄	NO	NO
EBC (3)		O ₃	O ₃
EBC (4)		HC (reactive)	HC1
EBC (5)		HNO ₂	HC2
EBC (6)			HC3
EBC (7)			HC4
EBC (8)			N ₂ O ₅
EBC (9)			HNO ₃
EBC (10)			HO ₂
EBC (11)			HO ₂
EBC (12)			H ₂ O ₂
EBC (13)			ROO
EBC (14)			RCOO
EBC (15)			

If IDOCDEM = 3, 6 or 15, at least one background concentration data must be specified for every K level; a maximum of 50 background concentration data sites may be specified.

4.3.12 Global Parameters Data

The global parameter data record specifies spatially invariant data required for conversion of emission g/sec to emissions ppm (the calculation of plume rise and parameters needed for photochemical calculations). If IDOCDEM = 0, this record is not required. Otherwise, this record is required each hour and concludes the hour input unit. The format of this record is as follows:

F5.0, ↑ TGRND	F5.0, ↑ SOLAR	F5.0, ↑ PRESR	F5.0, ↑ AMBCO	F5.0 ↑ AMHZO
---------------------	---------------------	---------------------	---------------------	--------------------

With the following global parameter definitions:

TGRND: air temperature at the surface °C (specify 20° as a default value)

SOLAR: ambient insolation in watts/m² (needed for photochemical simulations)

PRESR: ambient atmospheric pressure in millibars (specify 1000. as a default value)

AMBCO: ambient carbon monoxide in ppm (required for EPA photochemistry, IDOCM = 15)

AMHZO: ambient water vapor concentration in ppm (required for EPA photochemistry, IDOCM = 15)

4.4 PROGRAM INPUT SUMMARY

Table 6 provides a summary of program input.

Table 6. Summary of Input Data File.

5. CAVEATS

This section provides some helpful points on assembling the data and interpreting the output.

5.1 STABILITY ASSIGNMENT

Since the stability profile (stored in STAB) alters the computed wind field it is important to assign the stability layer by layer such that the intended profile is effected. The point to remember is that the stability of a given layer is assigned to the top interface of that layer. For example, if it is desired to simulate an atmosphere with the third layer stable and all the other layers unstable, the following assignment must be followed:

physically	assignment
A	A
A	A
A	A
A	A
A	A
E	E
A	E
A	A

thus STAB (2) = 5
and STAB (3) = 5

5.2 GRID SIZE

If IMPACT is used to evaluate the effects on air quality due to point sources, the grid size (DX, DY) should be equal to or less than the plume width at the surface concentration maximum. If DX or DY is too large, the maximum surface concentration will be smoothed and lower concentration values will be predicted. Typical zone sizes are from 500 to 1000 meters for DX, DY and 25 to 100 meters for DZ.

5.3 NUMBER OF SPECIFIED WIND SITES

If an excessive number of wind sites are used, the assignment of high confidence levels to each site will result in an unrealistic wind field if the data is not compatable. A rule of thumb for determining the maximum number of high confidence level wind sites is $N = (NX \cdot NY) / 40$. If the number of wind sites exceeds N, reduce the confidence values assigned to the wind data.

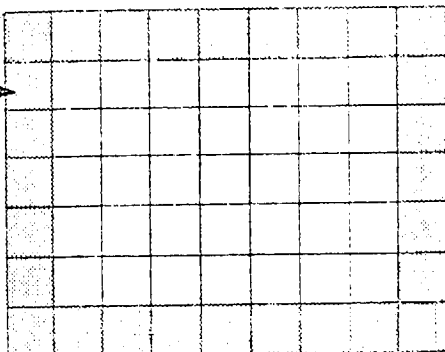
5.4 MYRUP/RANZIERI DIFFUSIVITY MODEL

When using the Myrup/Ranzieri Model (IDODIF = 1), the user should take care that an inversion base and top are defined everywhere in the grid. That is, the stability values go from less than 4.5 at the surface to greater than 4.5 and then to less than 4.5 again. Otherwise, the model may calculate large vertical diffusivities (limited to 1000 m²/sec) even under stable conditions when DZ·NZ is greater than 500 to 1000 meters. The advection/diffusion time step will be limited in these circumstances to approximately $\Delta t = 0.0008 (DZ)^2$; if DZ is equal to 100 meters, then Δt equal 8 seconds. A time step of 8 seconds requires 450 advection/diffusion cycles per hour which will result in long run times for multiple-species multiple-hour simulations.

5.5 LOCATION OF POINT AND AREA SOURCES

Due to the boundary conditions specified for the second order advection/diffusion algorithm, placing point or area sources in cells at the edge of the computational domain will result in an over prediction of pollutant concentrations. The user should take care that all point and area sources are at least one cell removed from the boundary.

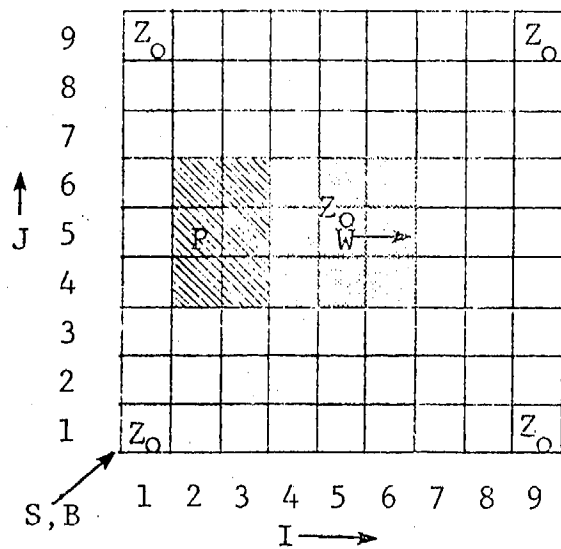
Avoid placing
sources in the
shaded cells.



6.0 EXAMPLE PROBLEM

A simplified exemplary problem designed to demonstrate several of the features and options available in the IMPACT code is described below. Section 6.1 illustrates the input file and section 6.2 is the program output.

The problem illustrated below is a grid of dimensions $NX = NY = 9$, $NZ = 7$; $DX = 750$, $DY = 500$, $DZ = 100$. The shaded area indicates the non-zero terrain (height = 1000 feet). Five surface roughness data are specified (Z_0), although they are not required since $IDODIF = 3$ (DEPICT diffusivity).



The problem is to be run for 2 hours, with wind and stability data specified (wind at w, stability at s) each hour. Sulfate chemistry is selected ($IDOCDEM = 3$) and there is simulation of both a point source, P, with a user-specified Δh and emission equal to 10 g/sec tracer, 500 g/sec SO_2 , and no SO_4 and an area source, indicated by the cross hatched area, with emissions per grid cell equal to 2 g/sec tracer and 270 g/sec SO_2 . Background concentrations (B) are set to 0.0 for all levels and all species. The wind direction remains constant at 270° for both hours, while the wind speed changes from 3.0 m/sec to 1.5 m/sec. The stability profile is unstable to neutral to stable the first hour, changing to completely stable the second hour.

6.1 EXAMPLE PROBLEM INPUT FILE

```
GRDIT Y M P A C T T E S I P R O R L E H
OPTION DX=750,DY=500,DX1=100,DX2=100,DY1=9,DY2=7, &END
OPTION IDOWNC=1,IDOCEM=3,IDOPLH=8,IDOUDIF=3, &END
OUTPUT IDOPLT=1,IDOCL=0,IDUPRN=1,IPSAGV=2,NUMHRS=2,IDSUR=1, &END
1 6 0. 0.1000.1000.1000. 0. 0.
1 5 0. 0.1000.1000.1000. 0. 0.
1 4 0. 0.1000.1000.1000. 0. 0.
*1
5 1 1 *1
1 9 1 *1
9 9 1 *1
9 5 1 *
5 5 1 *
1 2 5 .8 HOUR 1
5 270 .
1 1 2
1 1 3 2. 3. 4. 5. 6.
2. 2. 3. 4. 5. 6. 7.
1 2 5 100. 0. 0. 0. 120.
10. 500. 0. 0. 0. 0. 0.
1 3 3 4 6
2. 270. 0.
1 1 3
1 1 1 0. 0. 0.
0. 0. 0. 0. 0.
0. 0. 0. 0. 0.
0. 0. 0. 0. 0.
0. 0. 0. 0. 0.
20. 0.1^00. 0. 0. 0.
1 3 5 1 *
5 5 1 *
1.5 270. 1.
1 1 3
1 1 1 0. 0. 0.
6. 6. 6. 6. 6.
25. 0.1^00. 0. 0. 0.
```

5.2 EXAMPLE PROBLEM OUTPUT

```
SPUDIT T-M PART TEST PROGRAM  
DX= 750.0000  NY= 500.0000  NX= 100.0000  Q, NYZ= Q, Y2= Q,  
$  
SOLUTION IDOMD= 1, INPUTME= 3, IDOP1 ME= 8, IDODF= 3, $  
IDOMD= 1, INPUTME= 1, INPUTT= 0, IDOPRQ= 1, IDOMAVG= 2.00000  
SOLUTION IDOMD= 1, INPUTME= 1, INPUTT= 1, IDOPRQ= 1, IDOMAVG= 2.00000  
IDOSUR= 1, S
```

DRAFT 1 - QUESTIONS CONCERNANT LES CIRCONNÉANCES

ALLAN FARRICK
PAULINE SKLAFFER
JAMES TAIT
JOHN WILLIAMS
(213) 449-0514

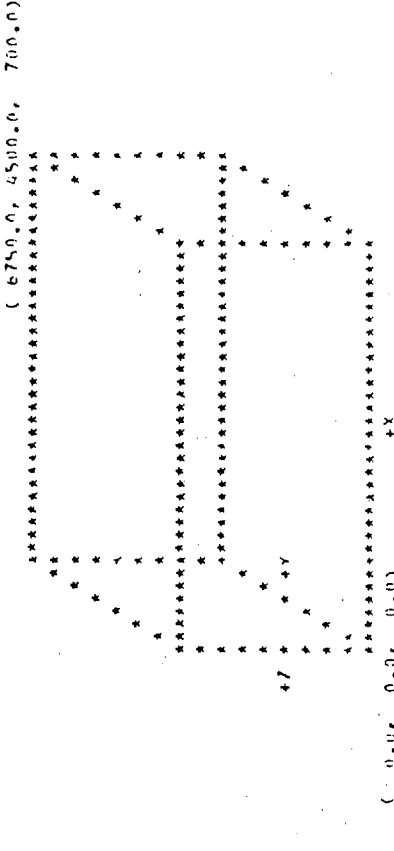
* INITIAL STATE OF THE SYSTEM *
* *****

IMPACT TEST PROFILE

GRID DEFINITION

THEK ARE 9 CELLS IN THE X DIRECTION WITH A DELTA X OF 750.0M YIELDING A TOTAL EXTENT OF 6750.0M
THERE ARE 9 CELLS IN THE Y DIRECTION WITH A DELTA Y OF 500.0M YIELDING A TOTAL EXTENT OF 4500.0M
THE ARE 7 CELLS IN THE Z DIRECTION WITH A DELTA Z OF 100.0M YIELDING A TOTAL EXTENT OF 700.0M

GRID INITIALIZATION



REQUESTED WINDS OF OPERATION

WIND FIELD

THE WIND FIELD IS TO BE GENERATED BY WEST

TURBULENCE

DIFFUSIVITIES TO BE DEVELOPED FROM DEPICT MEDIUM

PLUME RISE

PLUME RISE GIVEN BY USE IN 'DH' FORM

CHIMISTRY

SPECIFIC CHIMISTRY HAS BEEN PROVIDED

TIMING PARAMETERS

RUN TIME

A 2 HOUR SIMULATION HAS BEEN REQUESTED.

OUTPUT OPTIONS

PRINTER PLOTS HAVE BEEN REQUESTED EVERY 1 HOURS
PRINTER PLOTS HAVE BEEN REQUESTED EVERY 1 HOURS
SURFACE CONCENTRATIONS ARE 2 HOUR AVERAGES

INPUT TERRAIN DATA

I	J	ELEVATIONS (FT)				
1	6	0.	0.	1000.	1000.	0.
1	5	0.	0.	1000.	1000.	0.
1	4	0.	0.	1000.	1000.	0.

REFERENCE HEIGHT AT BOTTOM OF GRID (M) = 0.00

SURFACE ROUGHNESS DATA

N	I	J	ZI (M)
1	1	1	0.10
2	1	9	0.10
3	9	1	0.10
4	9	9	0.10
5	5	5	1.00

TERRAIN HEIGHT CONTOURS (M)

CONTOUR LEVELS

PLOT CHAR INTERVAL

	0.00000E+00	1.52402E+01	3.04804E+01	4.57206E+01	6.09608E+01	7.62010E+01	9.14412E+01	1.06681E+02	1.21922E+02	1.37162E+02	1.52402E+02	1.67642E+02	1.82882E+02	1.98123E+02	2.13363E+02	2.28603E+02	2.43843E+02	2.59083E+02	2.74323E+02	2.89563E+02	3.04803E+02
0	0.00000E+00	1.52402E+01	3.04804E+01	4.57206E+01	6.09608E+01	7.62010E+01	9.14412E+01	1.06681E+02	1.21922E+02	1.37162E+02	1.52402E+02	1.67642E+02	1.82882E+02	1.98123E+02	2.13363E+02	2.28603E+02	2.43843E+02	2.59083E+02	2.74323E+02	2.89563E+02	3.04803E+02
1	1.52402E+01	3.04804E+01	4.57206E+01	6.09608E+01	7.62010E+01	9.14412E+01	1.06681E+02	1.21922E+02	1.37162E+02	1.52402E+02	1.67642E+02	1.82882E+02	1.98123E+02	2.13363E+02	2.28603E+02	2.43843E+02	2.59083E+02	2.74323E+02	2.89563E+02	3.04803E+02	
2	3.09604E+01	4.62006E+01	6.14408E+01	7.66810E+01	9.19212E+01	1.07214E+02	1.22516E+02	1.37818E+02	1.53120E+02	1.68422E+02	1.83724E+02	1.99026E+02	2.14328E+02	2.29630E+02	2.44932E+02	2.60234E+02	2.75536E+02	2.90838E+02	3.06140E+02		
3	4.62006E+01	6.14408E+01	7.66810E+01	9.19212E+01	1.07214E+02	1.22516E+02	1.37818E+02	1.53120E+02	1.68422E+02	1.83724E+02	1.99026E+02	2.14328E+02	2.29630E+02	2.44932E+02	2.60234E+02	2.75536E+02	2.90838E+02	3.06140E+02			
4	6.14408E+01	7.66810E+01	9.19212E+01	1.07214E+02	1.22516E+02	1.37818E+02	1.53120E+02	1.68422E+02	1.83724E+02	1.99026E+02	2.14328E+02	2.29630E+02	2.44932E+02	2.60234E+02	2.75536E+02	2.90838E+02	3.06140E+02				
5	7.66810E+01	9.19212E+01	1.07214E+02	1.22516E+02	1.37818E+02	1.53120E+02	1.68422E+02	1.83724E+02	1.99026E+02	2.14328E+02	2.29630E+02	2.44932E+02	2.60234E+02	2.75536E+02	2.90838E+02	3.06140E+02					
6	9.19212E+01	1.07214E+02	1.22516E+02	1.37818E+02	1.53120E+02	1.68422E+02	1.83724E+02	1.99026E+02	2.14328E+02	2.29630E+02	2.44932E+02	2.60234E+02	2.75536E+02	2.90838E+02	3.06140E+02						
7	1.07214E+02	1.22516E+02	1.37818E+02	1.53120E+02	1.68422E+02	1.83724E+02	1.99026E+02	2.14328E+02	2.29630E+02	2.44932E+02	2.60234E+02	2.75536E+02	2.90838E+02	3.06140E+02							
8	1.22516E+02	1.37818E+02	1.53120E+02	1.68422E+02	1.83724E+02	1.99026E+02	2.14328E+02	2.29630E+02	2.44932E+02	2.60234E+02	2.75536E+02	2.90838E+02	3.06140E+02								
9	1.37818E+02	1.53120E+02	1.68422E+02	1.83724E+02	1.99026E+02	2.14328E+02	2.29630E+02	2.44932E+02	2.60234E+02	2.75536E+02	2.90838E+02	3.06140E+02									

TERRAIN HEIGHT PRINT (M)

MULTIPLIFER= 1.00000E+01

I= 1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22
J= 9	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
J= 8	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
J= 7	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
J= 6	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
J= 5	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
J= 4	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
J= 3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
J= 2	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
J= 1	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0

CELLULARIZED TERRAIN HEIGHT

I=	1	2	3	4	5	6	7	8	9
J=	9	0	0	0	0	0	0	0	0
J=	8	0	0	0	0	0	0	0	0
J=	7	0	0	0	0	0	0	0	0
J=	6	0	0	0	3	3	3	3	3
J=	5	0	0	0	3	3	3	3	3
J=	4	0	0	0	3	3	3	3	3
J=	3	0	0	0	0	0	0	0	0
J=	2	0	0	0	0	0	0	0	0
J=	1	0	0	0	0	0	0	0	0

SURFACE BRIGHTNESS FORTUNES (4)

CAPTION LEVELS

PLAT CHAO

	INTERVAL	VAL
0	1.0000E+01	1.0000E+01
	1.5000E+01	1.0000E+01
1	1.9000E+01	2.3500E+01
	2.3500E+01	2.36043E+01
2	2.8000E+01	3.2500E+01
	3.2500E+01	3.7600E+01
3	3.7500E+01	4.1500E+01
	4.1500E+01	4.6900E+01
4	4.6500E+01	5.0500E+01
	5.0500E+01	5.4500E+01
5	5.5000E+01	5.8500E+01
	5.8500E+01	6.2500E+01
6	6.3500E+01	6.7500E+01
	6.7500E+01	7.1500E+01
7	7.3000E+01	7.7500E+01
	7.7500E+01	8.2000E+01
8	8.2500E+01	8.7500E+01
	8.7500E+01	9.2500E+01
9	9.3000E+01	9.8000E+01
	9.8000E+01	1.0300E+02

***** VAL = 1.0000E+00

***** VAL = 1.0000E+01

SURFACE BRIGHTNESS OBTAIN

MULTIPLIER = 1.0000E+01

I=	1	2	3	4	5	6	7	8	9
J= 9	1	1	1	1	1	1	1	1	1
J= 8	1	2	1	2	1	2	1	2	1
J= 7	2	3	5	7	6	8	5	7	2
J= 6	2	3	5	8	2	5	3	2	1
J= 5	3	4	6	9	10	6	4	4	4
J= 4	2	3	4	5	9	6	3	2	1
J= 3	2	3	4	7	6	7	6	3	2
J= 2	1	2	3	4	6	4	2	1	1
J= 1	1	1	3	4	4	3	1	1	1

THE PRACTICAL STUDY OF THE

SIMILARITY OF CLOUDS IN DATA

25

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Age	Sex	Mean	SD	N
18-29	Male	1.00	0.00	10
18-29	Female	1.00	0.00	10
30-39	Male	1.00	0.00	10
30-39	Female	1.00	0.00	10
40-49	Male	1.00	0.00	10
40-49	Female	1.00	0.00	10
50-59	Male	1.00	0.00	10
50-59	Female	1.00	0.00	10
60-69	Male	1.00	0.00	10
60-69	Female	1.00	0.00	10
70+	Male	1.00	0.00	10
70+	Female	1.00	0.00	10

* CIRCUIT SIMULATION TIME 10 HRS

ESTATE PLANNING DATA

CHART 1
200
200
200
500
1000
1000

WIND AND PRESSURE PROFILE

* SUMMARY OF CURRENT EMISSIONS DATA
* *****

POINT SOURCES (GM/S)

N	J	EMISSIONS	STACK HEIGHT (M)	STACK TEMP (C)	FLOW RATE (M3/S)	STACK DIA (M)	EMISSION (M3/S)
1	S	SF6	100.0	0.0	0.0	0.0	0.0
1	S	SO2	500.0	0.0	0.0	0.0	0.0
1	S	SiH4	10.0	0.0	0.0	0.0	0.0

APFA SOURCES (GM/S)

N	IL	1H	JH	SF6	SiH4	SO2
1	2	3	4	6	7	0.0

BACKGROUND CONCENTRATION LEVELS (PPM)

N	1	2	3	4	5	6	7
1	1	1	1	1	0.00	0.00	0.00
2					1.00	1.00	1.00
3					0.00	0.00	0.00
4					0.00	0.00	0.00
5					0.00	0.00	0.00
6					0.00	0.00	0.00
7					0.00	0.00	0.00

GLOBAL ATMOSPHERIC PARAMETERS

GROUND TEMPERATURE	= 2.9000E+11 (DEG C)
SOLAR INSULATION	= 0.0000E+00 (W/M2)
SURFACE PRESSURE	= 1.0000E+05 (PA)
AMBIENT CO2	= 0.0000E+00 (PPM)
AMBIENT O2	= 0.0000E+00 (PPM)

BOUNDARY FLUX ADJUSTMENT

FLUX IN	1.740×10^{-6}
FLUX OUT	1.740×10^{-6}
FAC1- _W A	1.000×10^{-6}

WIND FIELD ADJUSTMENT

MAXIMUM RESIDUAL DIVERGENCE = 2.460×10^{-6} AT CELL I = 6 J = 5 K = 6
 CELL VELOCITIES I = 5.224×10^{-6} I+1 = 7.304×10^{-6} I+2 = 2.054×10^{-6} K = -6.083×10^{-6}
 PERCENT ERROR IN VELOCITY I = 3.648×10^{-6} Y = 1.904×10^{-6} Z = 1.247×10^{-6}

POINT SOURCE PLUME RISE

N	I	J	K	PLUME RISE	STKE HGT	TERRALN HGT	WEIN SITTR	THMOP (SRA)	THMOP (SRA)
1	2	5	2	1.20×10^{-6}	1.00×10^{-6}	0.00×10^{-6}	0.00×10^{-6}	0.00×10^{-6}	0.00×10^{-6}

POLLUTION ADVECTION DIFFUSION TIME STEP

DI	STEPS/AD	X-DI	Y-DI	Z-DI	ALPHA
52.94	68	7.70	1.08.40	5.6.40	3.95

SURFACE PLOT OF U VELOCITY

CENTROID LEVELS

PLOT CHAR INTERVAL

	0	1	2	3	4	5	6	7	8	9
0	0.0000E+00	-1.71419E+01	-1.71419E+01	-3.4285E+01	-5.14256E+01	-6.85575E+01	-8.57093E+01	-1.02851E+01	-1.19999E+01	-1.37153E+01
1	1.71419E+01	-3.4285E+01	-5.14256E+01	-6.85575E+01	-8.57093E+01	-1.02851E+01	-1.19999E+01	-1.37153E+01	-1.54277E+01	-1.71419E+01
2	3.42837E+01	-6.85575E+01	-8.57093E+01	-1.02851E+01	-1.19999E+01	-1.37153E+01	-1.54277E+01	-1.71419E+01	-1.88561E+01	-2.05702E+01
3	5.14256E+01	-8.57093E+01	-1.02851E+01	-1.19999E+01	-1.37153E+01	-1.54277E+01	-1.71419E+01	-1.88561E+01	-2.05702E+01	-2.22840E+01
4	6.85575E+01	-1.02851E+01	-1.19999E+01	-1.37153E+01	-1.54277E+01	-1.71419E+01	-1.88561E+01	-2.05702E+01	-2.22840E+01	-2.40926E+01
5	8.57093E+01	-1.19999E+01	-1.37153E+01	-1.54277E+01	-1.71419E+01	-1.88561E+01	-2.05702E+01	-2.22840E+01	-2.40926E+01	-2.58712E+01
6	1.02851E+01	-1.37153E+01	-1.54277E+01	-1.71419E+01	-1.88561E+01	-2.05702E+01	-2.22840E+01	-2.40926E+01	-2.58712E+01	-2.74271E+01
7	1.19999E+01	-1.54277E+01	-1.71419E+01	-1.88561E+01	-2.05702E+01	-2.22840E+01	-2.40926E+01	-2.58712E+01	-2.74271E+01	-2.91412E+01
8	1.37153E+01	-1.71419E+01	-1.88561E+01	-2.05702E+01	-2.22840E+01	-2.40926E+01	-2.58712E+01	-2.74271E+01	-2.91412E+01	-3.08554E+01
9	1.54277E+01	-1.88561E+01	-2.05702E+01	-2.22840E+01	-2.40926E+01	-2.58712E+01	-2.74271E+01	-2.91412E+01	-3.08554E+01	-3.25695E+01

SURFACE PLOT OF V VELOCITY

CENTROID LEVELS

PLOT CHAR INTERVAL

	0	1	2	3	4	5	6	7	8	9
0	-6.01677E-01	-6.13509E-01	-5.45341E-01	-4.09005E-01	-4.77174E-01	-5.55555E-01	-5.55555E-01	-5.55555E-01	-5.55555E-01	-5.55555E-01
1	-6.13509E-01	-5.45341E-01	-4.09005E-01	-4.77174E-01	-5.55555E-01	-5.55555E-01	-5.55555E-01	-5.55555E-01	-5.55555E-01	-5.55555E-01
2	-5.45341E-01	-4.09005E-01	-4.77174E-01	-5.55555E-01						
3	-4.09005E-01	-4.77174E-01	-5.55555E-01							
4	-4.77174E-01	-5.55555E-01								
5	-5.55555E-01									
6	-6.81794E-02	-1.36333E-01								
7	-2.04507E-01									
8	-2.72575E-01	-3.40843E-01								
9	-3.40843E-01	-4.09014E-01								

SURFACE CHEMISTRIES

卷之三

INTRODUCTION

卷之三

卷之三

ECONOMIC GROWTH AND INFLATION

卷之三

$$60 + 100 \cdot 5^2 = 2500 - 2500 = 0$$

2.2500000000000002

$$2.50000E+00 = 2.50000 \times 10^0$$

0	2-05000F+00	=	2-05000F+00						
1	2-05000F+01	-	2-10000F+01	*	00000000000000000000000000000000*				
2	2-10000F+00	-	2-15000F+00	*	00000000000000000000000000000000*				
3	2-15000F+00	-	2-20000F+00	*	00000000000000000000000000000000*				
4	2-20000F+00	-	2-25000F+00	*	00000000000000000000000000000000*				
5	2-25000F+00	-	2-30000F+00	*	00000000000000000000000000000000*				
6	2-30000F+00	-	2-35000F+00	*	00000000000000000000000000000000*				
7	2-35000F+00	-	2-40000F+00	*	00000000000000000000000000000000*				
8	2-40000F+00	-	2-45000F+00	*	00000000000000000000000000000000*				
9	2-45000F+00	-	2-50000F+00	*	00000000000000000000000000000000*				

SUGAR-FREE PLATE FOR A DIFFERENTIALLY

卷之三

卷之三

2.09171F+01 = 2.09171E+01

SURFACE POINTS & PRECISION

COMMON LEVELS

PLDT CHAR INTERVAL

0	0.0000E+00 = 1.20390E+00
1	1.20390E+00 = 2.09171E+00
2	2.09171E+00 = 3.65117E+00
3	3.65117E+00 = 4.61160E+00
4	4.61160E+00 = 5.61200E+00
5	5.61200E+00 = 6.61240E+00
6	6.61240E+00 = 7.22540E+00
7	7.22540E+00 = 7.82730E+00
8	7.82730E+00 = 8.42130E+00
9	8.42130E+00 = 9.03120E+00
10	9.03120E+00 = 9.63120E+00
11	9.63120E+00 = 1.02350E+01
12	1.02350E+01 = 1.08350E+01
13	1.08350E+01 = 1.20390E+01
14	1.20390E+01 = 1.32420E+01
15	1.32420E+01 = 1.44460E+01
16	1.44460E+01 = 1.56500E+01
17	1.56500E+01 = 1.68540E+01
18	1.68540E+01 = 1.80580E+01
19	1.80580E+01 = 1.92620E+01
20	1.92620E+01 = 2.04660E+01
21	2.04660E+01 = 2.16700E+01
22	2.16700E+01 = 2.28740E+01
23	2.28740E+01 = 2.40780E+01
24	2.40780E+01 = 2.52820E+01
25	2.52820E+01 = 2.64860E+01
26	2.64860E+01 = 2.76900E+01
27	2.76900E+01 = 2.88940E+01
28	2.88940E+01 = 3.00980E+01
29	3.00980E+01 = 3.12920E+01
30	3.12920E+01 = 3.24960E+01
31	3.24960E+01 = 3.36900E+01
32	3.36900E+01 = 3.48940E+01
33	3.48940E+01 = 3.60980E+01
34	3.60980E+01 = 3.73020E+01
35	3.73020E+01 = 3.85060E+01
36	3.85060E+01 = 3.97100E+01
37	3.97100E+01 = 4.09140E+01
38	4.09140E+01 = 4.21180E+01
39	4.21180E+01 = 4.33220E+01
40	4.33220E+01 = 4.45260E+01
41	4.45260E+01 = 4.57300E+01
42	4.57300E+01 = 4.69340E+01
43	4.69340E+01 = 4.81380E+01
44	4.81380E+01 = 4.93420E+01
45	4.93420E+01 = 5.05460E+01
46	5.05460E+01 = 5.17500E+01
47	5.17500E+01 = 5.29540E+01
48	5.29540E+01 = 5.41580E+01
49	5.41580E+01 = 5.53620E+01
50	5.53620E+01 = 5.65660E+01
51	5.65660E+01 = 5.77700E+01
52	5.77700E+01 = 5.89740E+01
53	5.89740E+01 = 6.01780E+01
54	6.01780E+01 = 6.13820E+01
55	6.13820E+01 = 6.25860E+01
56	6.25860E+01 = 6.37900E+01
57	6.37900E+01 = 6.49940E+01
58	6.49940E+01 = 6.61980E+01
59	6.61980E+01 = 6.74020E+01
60	6.74020E+01 = 6.86060E+01
61	6.86060E+01 = 6.98100E+01
62	6.98100E+01 = 7.10140E+01
63	7.10140E+01 = 7.22180E+01
64	7.22180E+01 = 7.34220E+01
65	7.34220E+01 = 7.46260E+01
66	7.46260E+01 = 7.58300E+01
67	7.58300E+01 = 7.70340E+01
68	7.70340E+01 = 7.82380E+01
69	7.82380E+01 = 7.94420E+01
70	7.94420E+01 = 8.06460E+01
71	8.06460E+01 = 8.18500E+01
72	8.18500E+01 = 8.30540E+01
73	8.30540E+01 = 8.42580E+01
74	8.42580E+01 = 8.54620E+01
75	8.54620E+01 = 8.66660E+01
76	8.66660E+01 = 8.78700E+01
77	8.78700E+01 = 8.90740E+01
78	8.90740E+01 = 9.02780E+01
79	9.02780E+01 = 9.14820E+01
80	9.14820E+01 = 9.26860E+01
81	9.26860E+01 = 9.38900E+01
82	9.38900E+01 = 9.50940E+01
83	9.50940E+01 = 9.62980E+01
84	9.62980E+01 = 9.75020E+01
85	9.75020E+01 = 9.87060E+01
86	9.87060E+01 = 9.99100E+01
87	9.99100E+01 = 1.01040E+02
88	1.01040E+02 = 1.02170E+02
89	1.02170E+02 = 1.03300E+02
90	1.03300E+02 = 1.04430E+02
91	1.04430E+02 = 1.05560E+02
92	1.05560E+02 = 1.06690E+02
93	1.06690E+02 = 1.07820E+02
94	1.07820E+02 = 1.08950E+02
95	1.08950E+02 = 1.10080E+02
96	1.10080E+02 = 1.11210E+02
97	1.11210E+02 = 1.12340E+02
98	1.12340E+02 = 1.13470E+02
99	1.13470E+02 = 1.14600E+02
100	1.14600E+02 = 1.15730E+02
101	1.15730E+02 = 1.16860E+02
102	1.16860E+02 = 1.17990E+02
103	1.17990E+02 = 1.19120E+02
104	1.19120E+02 = 1.20250E+02
105	1.20250E+02 = 1.21380E+02
106	1.21380E+02 = 1.22510E+02
107	1.22510E+02 = 1.23640E+02
108	1.23640E+02 = 1.24770E+02
109	1.24770E+02 = 1.25900E+02
110	1.25900E+02 = 1.27030E+02
111	1.27030E+02 = 1.28160E+02
112	1.28160E+02 = 1.29290E+02
113	1.29290E+02 = 1.30420E+02
114	1.30420E+02 = 1.31550E+02
115	1.31550E+02 = 1.32680E+02
116	1.32680E+02 = 1.33810E+02
117	1.33810E+02 = 1.34940E+02
118	1.34940E+02 = 1.36070E+02
119	1.36070E+02 = 1.37200E+02
120	1.37200E+02 = 1.38330E+02
121	1.38330E+02 = 1.39460E+02
122	1.39460E+02 = 1.40590E+02
123	1.40590E+02 = 1.41720E+02
124	1.41720E+02 = 1.42850E+02
125	1.42850E+02 = 1.43980E+02
126	1.43980E+02 = 1.45110E+02
127	1.45110E+02 = 1.46240E+02
128	1.46240E+02 = 1.47370E+02
129	1.47370E+02 = 1.48500E+02
130	1.48500E+02 = 1.49630E+02
131	1.49630E+02 = 1.50760E+02
132	1.50760E+02 = 1.51890E+02
133	1.51890E+02 = 1.53020E+02
134	1.53020E+02 = 1.54150E+02
135	1.54150E+02 = 1.55280E+02
136	1.55280E+02 = 1.56410E+02
137	1.56410E+02 = 1.57540E+02
138	1.57540E+02 = 1.58670E+02
139	1.58670E+02 = 1.59800E+02
140	1.59800E+02 = 1.60930E+02
141	1.60930E+02 = 1.62060E+02
142	1.62060E+02 = 1.63190E+02
143	1.63190E+02 = 1.64320E+02
144	1.64320E+02 = 1.65450E+02
145	1.65450E+02 = 1.66580E+02
146	1.66580E+02 = 1.67710E+02
147	1.67710E+02 = 1.68840E+02
148	1.68840E+02 = 1.70970E+02
149	1.70970E+02 = 1.73100E+02
150	1.73100E+02 = 1.76230E+02
151	1.76230E+02 = 1.79360E+02
152	1.79360E+02 = 1.82490E+02
153	1.82490E+02 = 1.85620E+02
154	1.85620E+02 = 1.88750E+02
155	1.88750E+02 = 1.91880E+02
156	1.91880E+02 = 1.95010E+02
157	1.95010E+02 = 1.98140E+02
158	1.98140E+02 = 2.01270E+02
159	2.01270E+02 = 2.04400E+02
160	2.04400E+02 = 2.07530E+02
161	2.07530E+02 = 2.10660E+02
162	2.10660E+02 = 2.13790E+02
163	2.13790E+02 = 2.16920E+02
164	2.16920E+02 = 2.20050E+02
165	2.20050E+02 = 2.23180E+02
166	2.23180E+02 = 2.26310E+02
167	2.26310E+02 = 2.29440E+02
168	2.29440E+02 = 2.32570E+02
169	2.32570E+02 = 2.35700E+02
170	2.35700E+02 = 2.38830E+02
171	2.38830E+02 = 2.41960E+02
172	2.41960E+02 = 2.45090E+02
173	2.45090E+02 = 2.48220E+02
174	2.48220E+02 = 2.51350E+02
175	2.51350E+02 = 2.54480E+02
176	2.54480E+02 = 2.57610E+02
177	2.57610E+02 = 2.60740E+02
178	2.60740E+02 = 2.63870E+02
179	2.63870E+02 = 2.66990E+02
180	2.66990E+02 = 2.70120E+02
181	2.70120E+02 = 2.73250E+02
182	2.73250E+02 = 2.76380E+02
183	2.76380E+02 = 2.79510E+02
184	2.79510E+02 = 2.82640E+02
185	2.82640E+02 = 2.85770E+02
186	2.85770E+02 = 2.88900E+02
187	2.88900E+02 = 2.92030E+02
188	2.92030E+02 = 2.95160E+02
189	2.95160E+02 = 2.98290E+02
190	2.98290E+02 = 3.01420E+02
191	3.01420E+02 = 3.04550E+02
192	3.04550E+02 = 3.07680E+02
193	3.07680E+02 = 3.10810E+02
194	3.10810E+02 = 3.13940E+02
195	3.13940E+02 = 3.17070E+02
196	3.17070E+02 = 3.20200E+02
197	3.20200E+02 = 3.23330E+02
198	3.23330E+02 = 3.26460E+02
199	3.26460E+02 = 3.29590E+02
200	3.29590E+02 = 3.32720E+02
201	3.32720E+02 = 3.35850E+02
202	3.35850E+02 = 3.38980E+02
203	3.38980E+02 = 3.42110E+02
204	3.42110E+02 = 3.45240E+02
205	3.45240E+02 = 3.48370E+02
206	3.48370E+02 = 3.51500E+02
207	3.51500E+02 = 3.54630E+02
208	3.54630E+02 = 3.57760E+02
209	3.57760E+02 = 3.60890E+02
210	3.60890E+02 = 3.64020E+02
211	3.64020E+02 = 3.67150E+02
212	3.67150E+02 = 3.70280E+02
213	3.70280E+02 = 3.73410E+02
214	3.73410E+02 = 3.76540E+02
215	3.76540E+02 = 3.79670E+02
216	3.79670E+02 = 3.82800E+02
217	3.82800E+02 = 3.85930E+02
218	3.85930E+02 = 3.89060E+02
219	3.89060E+02 = 3.92190E+02
220	3.92190E+02 = 3.95320E+02
221	3.95320E+02 = 3.98450E+02
222	3.98450E+02 = 4.01580E+02
223	4.01580E+02 = 4.04710E+02
224	4.04710E+02 = 4.07840E+02
225	4.07840E+02 = 4.10970E+02
226	4.10970E+02 = 4.14100E+02
227	4.14100E+02 = 4.17230E+02
228	4.17230E+02 = 4.20360E+02
229	4.20360E+02 = 4.23490E+02
230	4.23490E+02 = 4.26620E+02
231	4.26620E+02 = 4.29750E+02
232	4.29750E+02 = 4.32880E+02
233	4.32880E+02 = 4.35010E+02
234	4.35010E+02 = 4.38140E+02
235	4.38140E+02 = 4.41270E+02
236	4.41270E+02 = 4.44400E+02
237	4.44400E+02 = 4.47530E+02
238	4.47530E+02 = 4.50660E+02
239	4.50660E+02 = 4.53790E+02
240	4.53790E+02 = 4.56920E+02
241	4.56920E+02 = 4.60050E+02
242	4.60050E+02 = 4.63180E+02
243	4.63180E+02 = 4.66310E+02
244	4.66310E+02 = 4.69440E+02
245	4.69440E+02 = 4.72570E+02
246	4.72570E+02 = 4.75700E+02
247	4.75700E+02 = 4.78830E+02
248	4.78830E+02 = 4.81960E+02
249	4.81960E+02 = 4.85090E+02
250	4.85090E+02 = 4.88220E+02
251	4.88220E+02 = 4.91350E+02
252	4.91350E+02 = 4.94480E+02
253	4.94480E+02 = 4.97610E+02
254	4.97610E+02 = 5.00740E+02
255	5.00740E+02 = 5.03870E+02
256	5.03870E+02 = 5.06990E+02
257	5.06990E+02 = 5.10120E+02
258	5.10120E+02 = 5.13250E+02
259	5.13250E+02 = 5.16380E+02
260	5.16380E+02 = 5.19510E+02
261	5.19510E+02 = 5.22640E+02
262	5.22640E+02 = 5.25770E+02
263	5.25770E+02 = 5.28900E+02
264	5.28900E+02 = 5.32030E+02
265	5.32030E+02 = 5.35160E+02
266	5.35160E+02 = 5.38290E+02
267	5.38290E+02 = 5.41420E+02
268	5.41420E+02 = 5.44550E+02
269	5.44550E+02 = 5.47680E+02

CONGNTRATION AT 11:00 1 HOURS AFTER START OF STIR, AT 1000' ELEVATION

SURFACE PLOT OF TRACER - UG/m**3

CONTOUR LEVELS

PLOT CHAR

INTERVAL

	0	1	2	3	4	5	6	7	8	9	0	
0	0.00000E+00	-	-	-	-	-	-	-	-	-	-	
1	1.59862E+00	-	3.17600E+00	-	4.76405E+00	-	6.35257E+00	-	7.94009E+00	-	9.52811E+00	-
2	-	6.75207E+01	-	7.94200E+01	-	9.52811E+01	-	1.11111E+02	-	1.27041E+02	-	
3	-	-	9.52811E+00	-	1.11111E+01	-	1.31111E+01	-	1.51111E+01	-	1.71111E+01	-
4	-	-	1.11111E+01	-	1.27041E+01	-	1.42922E+01	-	1.58962E+01	-	1.74882E+01	-
5	-	-	1.58962E+01	-	1.74882E+01	-	1.90542E+01	-	2.06442E+01	-	2.22323E+01	-
6	-	-	2.06442E+01	-	2.22323E+01	-	2.38205E+01	-	2.54085E+01	-	2.70963E+01	-
7	-	-	2.22323E+01	-	2.38205E+01	-	2.54085E+01	-	2.70963E+01	-	2.87843E+01	-
8	-	-	2.54085E+01	-	2.70963E+01	-	2.87843E+01	-	3.04723E+01	-	3.21604E+01	-
9	-	-	2.70963E+01	-	2.87843E+01	-	3.04723E+01	-	3.21604E+01	-	3.38484E+01	-
0	-	-	3.04723E+01	-	3.21604E+01	-	3.38484E+01	-	3.55364E+01	-	3.72245E+01	-
1	-	-	3.21604E+01	-	3.38484E+01	-	3.55364E+01	-	3.72245E+01	-	3.89125E+01	-
2	-	-	3.38484E+01	-	3.55364E+01	-	3.72245E+01	-	3.89125E+01	-	4.05995E+01	-
3	-	-	3.55364E+01	-	3.72245E+01	-	3.89125E+01	-	4.05995E+01	-	4.22875E+01	-
4	-	-	3.72245E+01	-	3.89125E+01	-	4.05995E+01	-	4.22875E+01	-	4.39755E+01	-
5	-	-	3.89125E+01	-	4.05995E+01	-	4.22875E+01	-	4.39755E+01	-	4.56635E+01	-
6	-	-	4.05995E+01	-	4.22875E+01	-	4.39755E+01	-	4.56635E+01	-	4.73515E+01	-
7	-	-	4.22875E+01	-	4.39755E+01	-	4.56635E+01	-	4.73515E+01	-	4.90395E+01	-
8	-	-	4.39755E+01	-	4.56635E+01	-	4.73515E+01	-	4.90395E+01	-	5.07275E+01	-
9	-	-	4.56635E+01	-	4.73515E+01	-	4.90395E+01	-	5.07275E+01	-	5.24155E+01	-

SURFACE PLOT OF SO₂ ppm

CONTOUR LEVELS

PLOT CHAR

	0	1	2	3	4	5	6	7	8	9	0	
0	0.00000E+00	-	-	-	-	-	-	-	-	-	-	
1	1.14443E+01	-	2.28887E+01	-	3.43329E+01	-	4.57772E+01	-	5.72215E+01	-	6.86658E+01	-
2	-	3.43329E+01	-	4.57772E+01	-	5.72215E+01	-	6.86658E+01	-	8.01101E+01	-	
3	-	-	4.57772E+01	-	5.72215E+01	-	6.86658E+01	-	8.01101E+01	-	9.15544E+01	-
4	-	-	5.72215E+01	-	6.86658E+01	-	8.01101E+01	-	9.15544E+01	-	1.02999E+02	-
5	-	-	6.86658E+01	-	8.01101E+01	-	9.15544E+01	-	1.02999E+02	-	1.14443E+02	-
6	-	-	8.01101E+01	-	9.15544E+01	-	1.02999E+02	-	1.14443E+02	-	1.25887E+02	-
7	-	-	9.15544E+01	-	1.02999E+02	-	1.14443E+02	-	1.25887E+02	-	1.37332E+02	-
8	-	-	1.02999E+02	-	1.14443E+02	-	1.25887E+02	-	1.37332E+02	-	1.48776E+02	-
9	-	-	1.14443E+02	-	1.25887E+02	-	1.37332E+02	-	1.48776E+02	-	1.60220E+02	-

SURFACE WELD		SO ₄ ppm	
PLATE CHAR	INTERVAL	CENTER LEVELS	CHAR
R	1.0-3109E+01	1.04553E+01	-
	1.04553E+01	2.05007E+01	-
0	2.05007E+01	2.07442E+01	-
	2.07442E+01	2.20885E+01	-
0	0.0-3000E+01	3.05491E+01	-
	3.05491E+01	6.09474E+01	-
1	6.09474E+01	9.16453E+01	-
	9.16453E+01	1.22140E+02	-
2	1.22140E+02	1.53742E+02	-
	1.53742E+02	1.85344E+02	-
3	1.85344E+02	2.16946E+02	-
	2.16946E+02	2.48548E+02	-
4	2.48548E+02	2.79150E+02	-
	2.79150E+02	3.09752E+02	-
5	3.09752E+02	3.41354E+02	-
	3.41354E+02	3.66461E+02	-
6	3.66461E+02	3.97130E+02	-
	3.97130E+02	4.28738E+02	-
7	4.28738E+02	4.59346E+02	-
	4.59346E+02	4.89954E+02	-
8	4.89954E+02	5.20562E+02	-
	5.20562E+02	5.51170E+02	-
9	5.51170E+02	5.81778E+02	-
	5.81778E+02	6.10967E+02	-

SURFACE PRINT OF VELICITY

MULTIPLIER= 1.00000E-01

I= 1	J= 1	2	3	4	5	6	7	8	9	10
J= 9	32	32	32	32	32	32	32	32	32	32
J= 8	34	34	34	34	34	34	34	34	34	34
J= 7	36	36	36	36	36	36	36	36	36	36
J= 6	30	29	29	29	29	29	29	29	29	29
J= 5	39	25	34	34	34	34	34	34	34	34
J= 4	30	29	25	34	34	34	34	34	34	34
J= 3	30	30	29	31	32	32	32	32	32	32
J= 2	30	30	30	30	30	30	30	30	30	30
J= 1	30	30	30	30	30	30	30	30	30	30

SURFACE PRINT OF VELICITY

MULTIPLIER= 1.00000E-02

I= 1	J= 10	9	8	7	6	5	4	3	2	1
J= 9	1	2	4	1	0	-1	-3	-5	-7	-9
J= 8	1	4	16	2	0	-3	-13	-3	-1	0
J= 7	1	8	68	0	0	-58	-6	-1	0	0
J= 6	0	2	13	1	0	-3	-11	-2	0	0
J= 5	0	-2	13	-1	0	5	11	2	0	0
J= 4	-1	-8	-65	-2	1	16	54	6	1	0
J= 3	-1	-6	-16	-2	0	3	15	3	1	0
J= 2	-1	-2	-4	-1	0	1	4	1	0	0
J= 1	0	-1	-2	-1	0	1	1	1	0	0

SURFACE PRINT OF STABILITY

MULTIPLIER= 1.00000E-03

I= 1	J= 9	20	20	20	20	20	20	20	20	20
J= 8	20	20	20	20	20	20	20	20	20	20
J= 7	20	20	20	20	20	20	20	20	20	20
J= 6	20	20	20	20	20	20	20	20	20	20
J= 5	20	20	20	20	20	20	20	20	20	20
J= 4	20	20	20	20	20	20	20	20	20	20
J= 3	20	20	20	20	20	20	20	20	20	20

ν^2
 J_2

SUPPLY CURRENT $\propto k \cdot 2^{(J_1+J_2)/2}$

MULTIPLIER = $1.72 \cdot 10^{-9} e^{J_1 + J_2}$

J_1	0	1	2	3	4	5	6	7	8	9	10
J_2	0	24	24	24	24	24	24	24	24	24	24
	1	24	24	24	24	24	24	24	24	24	24
2		24	24	24	24	24	24	24	24	24	24
3			24	24	24	24	24	24	24	24	24
4				24	24	24	24	24	24	24	24
5					24	24	24	24	24	24	24
6						24	24	24	24	24	24
7							24	24	24	24	24
8								24	24	24	24
9									24	24	24
10										24	24

SURFACE POWER OF DIFFUSIVITY

MULTIPLIER = $1.09 \cdot 10^{-14.69}$

J_1	0	1	2	3	4	5	6	7	8	9	10
J_2	10	24	24	24	24	24	24	24	24	24	24
	9	24	24	24	24	24	24	24	24	24	24
8		24	24	24	24	24	24	24	24	24	24
7			24	24	24	24	24	24	24	24	24
6				24	24	24	24	24	24	24	24
5					24	24	24	24	24	24	24
4						24	24	24	24	24	24
3							24	24	24	24	24
2								24	24	24	24
1									24	24	24
0										24	24

CONCENTRATION AT TIME 1 HOURS AFTER START OF CALCULATION

SURFACE PRINT OF PLACE 2 (S/MKS)

MULTIPLIER= 1.00000E+00

I= 1	2	3	4	5	6	7	8	9
J= 0	0	0	0	0	0	0	0	0
J= 1	0	0	0	0	0	0	0	0
J= 2	0	0	0	0	0	0	0	0
J= 3	0	0	0	0	0	0	0	0
J= 4	0	0	0	0	0	0	0	0
J= 5	0	0	0	0	0	0	0	0
J= 6	0	0	0	0	0	0	0	0
J= 7	0	0	0	0	0	0	0	0
J= 8	0	0	0	0	0	0	0	0
J= 9	0	0	0	0	0	0	0	0

SURFACE PRINT OF SO₂ ppm

MULTIPLIER= 1.00000E-02

I= 1	2	3	4	5	6	7	8	9
J= 0	0	0	0	0	0	0	0	0
J= 1	0	0	0	0	0	0	0	0
J= 2	0	0	0	0	0	0	0	0
J= 3	0	0	0	0	0	0	0	0
J= 4	0	0	0	0	0	0	0	0
J= 5	0	0	0	0	0	0	0	0
J= 6	0	0	0	0	0	0	0	0
J= 7	0	0	0	0	0	0	0	0
J= 8	0	0	0	0	0	0	0	0
J= 9	0	0	0	0	0	0	0	0

SURFACE PRINT OF SO₄ ppm

MULTIPLIER= 1.00000E-04

I= 1	2	3	4	5	6	7	8	9
J= 0	0	0	0	0	0	0	0	0
J= 1	0	0	0	0	0	0	0	0
J= 2	0	0	0	0	0	0	0	0
J= 3	0	0	0	0	0	0	0	0
J= 4	0	0	0	0	0	0	0	0
J= 5	0	0	0	0	0	0	0	0
J= 6	0	0	0	0	0	0	0	0
J= 7	0	0	0	0	0	0	0	0
J= 8	0	0	0	0	0	0	0	0
J= 9	0	0	0	0	0	0	0	0

6 12 3 9 0
6 3 4 0 0
4 12 2 0 0
5 12 0 0 0
8 3 0 0 0
6 3 0 0 0
6 12 2 0 0
6 12 2 0 0
6 9 0 0 0
6 9 0 0 0
6 9 0 0 0
6 9 0 0 0
4 3 2 1
J# J# J#

COMPARATIVE POSITION

```
*****
* SURFACE CURRENT DATA *
* *****
```

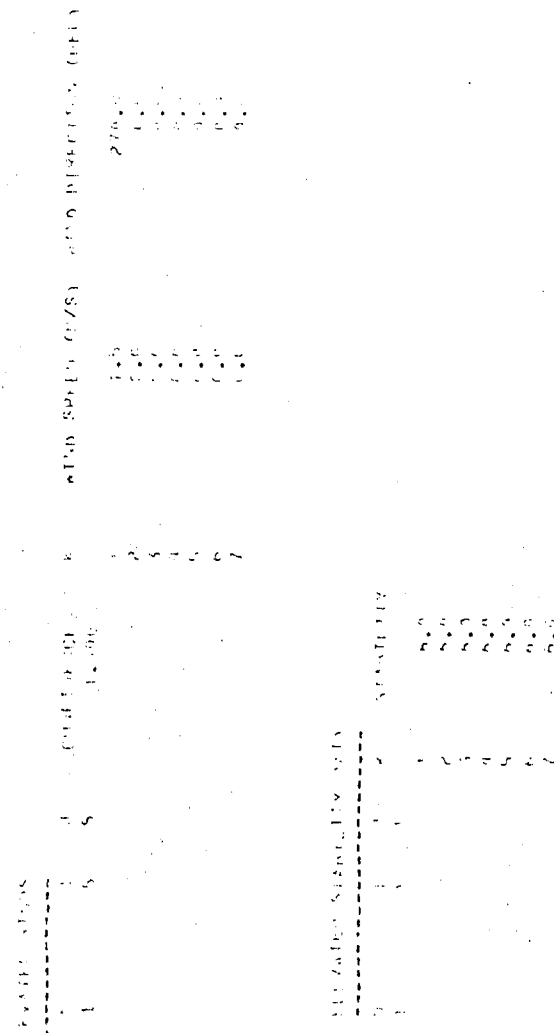


Fig. 1. Current Position

```
FIGURE 1. COMPARATIVE POSITION  
SURFACE CURRENT DATA  
SHEAR AND SWELL  
ADDITIONAL  
ADDITIONAL
```

BOUNDARY FLUX ADJUSTMENT

FLUX U_4	$1.1582E+07$
FLUX U_{11}	$1.1542E+07$
FACTOR κ	$1.0000E+00$

WIND FIELD ADJUSTMENT

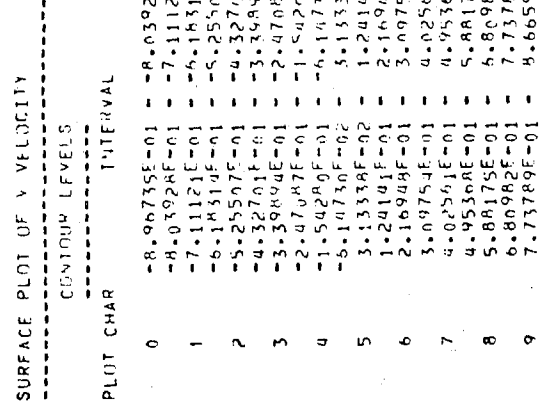
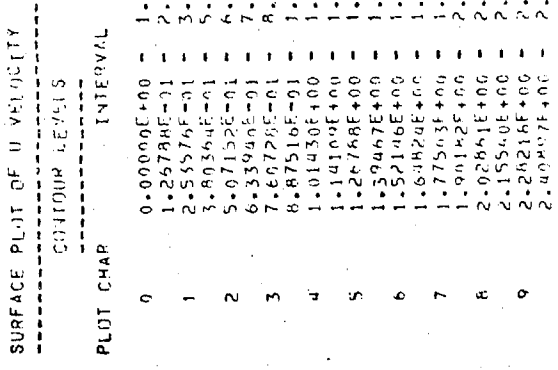
MAXIMUM RESIDUAL CONVERGENCE	$4.1755E-05$	AT CELL $J = 4, I = 4$
CELL VELocities	$1.5255E+00$	$J = 4, I = 4, \Delta U = 3.0499E-01$
PERCENT ERROR IN VELOCITY	$U = 1.675AE-02\%$	$V = 1.5415E-01\%$

POINT SOURCE PLUME ATSF

N	I	J	K	PLUMA X TSF	PLUMA Y TSF	PLUMA Z TSF	TSX TSF	TSY TSF	TSZ TSF	TSX TSF	TSY TSF	TSZ TSF
1	2	3	4	1.20	1.00	0.80	0.60	0.40	0.20	0.00	0.00	0.00

POLLUTANT ADVECTION DIFFUSION TIME STEP

DT	STEPS/HR	X-HR	Y-HR	Z-HR	ALPHA
12A.57	26	138.19	277.41	3153.22	0.05



$$A_{\text{min}} = 0.594 \times 10^{-1} = 0.59405 \times 10^{-1}$$

SURFACE PLATE OF STABILITY

CONTINUOUS EQUILIBRIUM

PLATE CURVE

	PLATE CURVE	TERMINAL
0	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
1	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
2	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
3	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
4	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
5	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
6	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
7	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
8	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
9	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$

$$\text{STABILISATION VALUE } V_{\text{st}} = 0.0000000000$$

$$\text{MAXIMUM VALUE } V_{\text{max}} = 0.0000000000$$

$$\text{MINIMUM VALUE } V_{\text{min}} = 0.0000000000$$

$$\text{STABILISATION VALUE } V_{\text{st}} = 0.0000000000$$

$$\text{MAXIMUM VALUE } V_{\text{max}} = 0.0000000000$$

$$\text{MINIMUM VALUE } V_{\text{min}} = 0.0000000000$$

SURFACE PLATE OF STABILITY

CONTINUOUS EQUILIBRIUM

PLATE CURVE

	PLATE CURVE	TERMINAL
0	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
1	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
2	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
3	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
4	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
5	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
6	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
7	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
8	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$
9	$b_{\text{min}} = 0.0000000000$	$b_{\text{max}} = 0.0000000000$

$$\text{STABILISATION VALUE } V_{\text{st}} = 0.0000000000$$

$$\text{MAXIMUM VALUE } V_{\text{max}} = 0.0000000000$$

$$\text{MINIMUM VALUE } V_{\text{min}} = 0.0000000000$$

CONCENTRATION AT 1 hr & 2 HOURS AFTER START OF SIMULATION

SURFACE PLOT OF TARGET SURFACE

CONTINUOUS LEVELS

PLANT CHAN

Plant Chan	0	1	2	3	4	5	6	7	P	Q	R	S
0	5.7419E+01											
1	1.1435E+01											
2	1.7225E+01	2.2067E+01										
3	3.4405E+01											
4	6.8818E+01											
5	1.1617E+01											
6	6.8818E+01											
7	2.0451E+01											
P	4.6167E+01											
Q	7.6154E+01											
R	1.2955E+02											
S	1.67937E+02											

SUPPLY POINT FOR SO₂ ppm

CONTINUOUS LEVELS

PLANT CHAN

Plant Chan	0	1	2	3	4	5	6	7	P	Q	R	S
0	1.0000E+00	2.0000E+00										
1	4.0215E+00											
2	7.5625E+00	9.3630E+00										
3	1.2146E+01	1.3768E+01										
4	1.9676E+01	2.2146E+01										
5	2.9013E+01	3.2705E+01										
6	4.9520E+01	5.4984E+01										
7	8.0898E+01	8.4450E+01										

B	3.9372E-01	-	4.18328E-01
	4.18328E-01	-	4.42935E-01
9	4.42935E-01	-	4.57543E-01
	4.57543E-01	-	4.92150E-01

SURFACE PLOT OF SO₄ ppm

COUNTUP LEVELS

PLOT CHAR VTFEVAL

0	0.00000E+00	-	1.60052E-01
	1.52190E-01	-	1.52190E-01
1	1.52190E-01	-	2.24286E-01
	2.24286E-01	-	3.01541E-01
2	3.01541E-01	-	3.80476E-01
	3.80476E-01	-	4.55511E-01
3	4.55511E-01	-	5.32657E-01
	5.32657E-01	-	6.18725E-01
4	6.18725E-01	-	6.94877E-01
	6.94877E-01	-	7.60952E-01
5	7.60952E-01	-	8.27017E-01
	8.27017E-01	-	8.84143E-01
6	8.84143E-01	-	9.39238E-01
	9.39238E-01	-	1.06555E-01
7	1.06555E-01	-	1.14145E-01
	1.14145E-01	-	1.21752E-01
8	1.21752E-01	-	1.29362E-01
	1.29362E-01	-	1.36911E-01
9	1.36911E-01	-	1.43591E-01
	1.43591E-01	-	1.52190E-01

2. HOUR AVERAGED CONCENTRATIONS

SURFACE PLOT OF TRACER 16/*****

CONTOUR LEVELS

PLOT CHAR T-INTERVAL

	0	1	2	3	4	5	6	7	8	9	10
0	0.00000E+000	-2.30217E+000	-4.60444E+000	-6.90652E+000	-9.20864E+000	-1.15107E+000	-1.38135E+000	-1.61152E+000	-1.84174E+000	-2.07196E+000	-2.30217E+000
1	4.00454E+000	-2.30217E+000	-4.60444E+000	-6.90652E+000	-9.20864E+000	-1.15107E+000	-1.38135E+000	-1.61152E+000	-1.84174E+000	-2.07196E+000	-2.30217E+000
2	6.90652E+000	-2.30217E+000	-4.60444E+000	-6.90652E+000	-9.20864E+000	-1.15107E+000	-1.38135E+000	-1.61152E+000	-1.84174E+000	-2.07196E+000	-2.30217E+000
3	1.15107E+000	-2.30217E+000	-4.60444E+000	-6.90652E+000	-9.20864E+000	-1.15107E+000	-1.38135E+000	-1.61152E+000	-1.84174E+000	-2.07196E+000	-2.30217E+000
4	1.61152E+000	-2.30217E+000	-4.60444E+000	-6.90652E+000	-9.20864E+000	-1.15107E+000	-1.38135E+000	-1.61152E+000	-1.84174E+000	-2.07196E+000	-2.30217E+000
5	1.84174E+000	-2.30217E+000	-4.60444E+000	-6.90652E+000	-9.20864E+000	-1.15107E+000	-1.38135E+000	-1.61152E+000	-1.84174E+000	-2.07196E+000	-2.30217E+000
6	2.07196E+000	-2.30217E+000	-4.60444E+000	-6.90652E+000	-9.20864E+000	-1.15107E+000	-1.38135E+000	-1.61152E+000	-1.84174E+000	-2.07196E+000	-2.30217E+000
7	3.22304E+001	-3.45323E+001	-3.68344E+001	-3.91364E+001	-4.14384E+001	-4.37404E+001	-4.60424E+001	-4.83444E+001	-5.06464E+001	-5.29484E+001	-5.52504E+001
8	3.45323E+001	-3.45323E+001	-3.68344E+001	-3.91364E+001	-4.14384E+001	-4.37404E+001	-4.60424E+001	-4.83444E+001	-5.06464E+001	-5.29484E+001	-5.52504E+001
9	4.14384E+001	-4.37404E+001	-4.60424E+001	-4.83444E+001	-5.06464E+001	-5.29484E+001	-5.52504E+001	-5.75524E+001	-5.98544E+001	-6.21564E+001	-6.44584E+001
10	4.37404E+001	-4.37404E+001	-4.60424E+001	-4.83444E+001	-5.06464E+001	-5.29484E+001	-5.52504E+001	-5.75524E+001	-5.98544E+001	-6.21564E+001	-6.44584E+001

SURFACE PLOT OF SO₂ ppm

CONTOUR LEVELS

PLOT CHAR T-INTERVAL

	0	1	2	3	4	5	6	7	8	9	10
0	0.00000E+000	-9.82843E-005	-1.96579E-002	-2.94868E-002	-3.93157E-002	-4.91446E-002	-5.89736E-002	-6.88025E-002	-7.86314E-002	-8.84604E-002	-9.82893E-002
1	9.82843E-005	-1.96579E-002	-2.94868E-002	-3.93157E-002	-4.91446E-002	-5.89736E-002	-6.88025E-002	-7.86314E-002	-8.84604E-002	-9.82893E-002	-1.08118E-001
2	1.96579E-002	-2.94868E-002	-3.93157E-002	-4.91446E-002	-5.89736E-002	-6.88025E-002	-7.86314E-002	-8.84604E-002	-9.82893E-002	-1.08118E-001	-1.17497E-001
3	2.94868E-002	-3.93157E-002	-4.91446E-002	-5.89736E-002	-6.88025E-002	-7.86314E-002	-8.84604E-002	-9.82893E-002	-1.08118E-001	-1.17497E-001	-1.27776E-001
4	3.93157E-002	-4.91446E-002	-5.89736E-002	-6.88025E-002	-7.86314E-002	-8.84604E-002	-9.82893E-002	-1.08118E-001	-1.17497E-001	-1.27776E-001	-1.37665E-001
5	4.91446E-002	-5.89736E-002	-6.88025E-002	-7.86314E-002	-8.84604E-002	-9.82893E-002	-1.08118E-001	-1.17497E-001	-1.27776E-001	-1.37665E-001	-1.47434E-001
6	5.89736E-002	-6.88025E-002	-7.86314E-002	-8.84604E-002	-9.82893E-002	-1.08118E-001	-1.17497E-001	-1.27776E-001	-1.37665E-001	-1.47434E-001	-1.57263E-001

8 $1.57263E-01$ = $1.57092E-01$
 1.57192E-01 = $1.570921E-01$
 9 $1.76921E-01$ = $1.76921E-01$
 1.86750E-01 = $1.86750E-01$
 1.26579E-01 =

SURFACE PLATE OF SO_4 ppm

CONTINUUM LEVELS

PLATE CHARGE INTERVAL

	0	$0.90000E+00$	$2.56289E-04$	$*00000000000000000000*$
		$2.56289E-04$	$5.12578E-04$	$*00000000000000000000*$
1	$5.12578E-04$	$7.68868E-04$	$*5.00055E-04$	$*00000000000000000000*$
	$7.68868E-04$	$1.02516E-03$	$*5.00055E-04$	$1.11111E-03$
2	$1.02516E-03$	$1.38445E-03$	$*00000000000000000000*$	$1.11111E-03$
	$1.38445E-03$	$1.53774E-03$	$*00000000000000000000*$	$1.11111E-03$
3	$1.53774E-03$	$1.79403E-03$	$*00000000000000000000*$	$1.11111E-03$
	$1.79403E-03$	$2.05031E-03$	$*00000000000000000000*$	$1.11111E-03$
4	$2.05031E-03$	$2.30660E-03$	$*00000000000000000000*$	$1.11111E-03$
	$2.30660E-03$	$2.56289E-03$	$*00000000000000000000*$	$1.11111E-03$
5	$2.56289E-03$	$2.81918E-03$	$*00000000000000000000*$	$1.11111E-03$
	$2.81918E-03$	$3.07547E-03$	$*00000000000000000000*$	$1.11111E-03$
6	$3.07547E-03$	$3.55517E-03$	$*00000000000000000000*$	$1.11111E-03$
	$3.55517E-03$	$3.83333E-03$	$*00000000000000000000*$	$1.11111E-03$
7	$3.83333E-03$	$4.10000E-03$	$*00000000000000000000*$	$1.11111E-03$
	$4.10000E-03$	$4.36667E-03$	$*00000000000000000000*$	$1.11111E-03$
8	$4.36667E-03$	$4.63333E-03$	$*00000000000000000000*$	$1.11111E-03$
	$4.63333E-03$	$4.90000E-03$	$*00000000000000000000*$	$1.11111E-03$
9	$4.90000E-03$	$5.16667E-03$	$*00000000000000000000*$	$1.11111E-03$
	$5.16667E-03$	$5.43333E-03$	$*00000000000000000000*$	$1.11111E-03$



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